Examples B-2150 through B-2173 are prepared from Scaffold C-32

		Janough B Z 170 a	о разралос		
Example#	R²	K,			
B-2150	F—			·	
B-2151	F—) F			
B-2152	F—	34			
B-2153	F—				
B-2154	F-	2/			
B-2155	F—{	2,1			
B-2156	F-{}	O BR			

					
Example#	R²	K,			
B-2157	F—				
B-2158	F—	0 - Z			
B-2159	F—	340			
B-2160	F—	3.4			
B-2161	F—				
B-2162	F—	-0,-			
B-2163	F—	F.r.			
B-2164	F—	N. T.			
B-2165	F—		D:		
B-2166	F—	7, 0 /0			

Example#	R²	К ₁		
B-2167	F—	7,8%		
B-2168	F—	7,500 F. 000		
B-2169	F—	Z NH		
B-2170	F—	Y		
B-2171	F—			
B-2172	F—	HN-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
B-2173	F—	PN 7		

Examples 2174 through B-2197 are prepared from Scaffold C-64

	Examples 2177	iniough 6-2197 are	P. 0 P	
Example#	R²	۴٦		
B-2174	F—			
B-2175	F—	° F		
B-2176	F—			
B-2177	F—			
B-2178	F	24		
B-2179	F-	2		
B-2180	F-) BR		

Example#	R²	В ₁		
B-2181	F—	£.		
B-2182	F—	Z 2-2		
B-2183	F-			
B-2184	F—	1, °-		
B-2185	F—	13		
B-2186	F-			
B-2187	F—	Gr.		
B-2188	F—	~~~.		
B-2189	F—			
B-2190	F-	7,0		

Example#	R²	₽,		
B-2191	F	78%		
B-2192		ξ		
B-2193	F—	LT O NH		
B-2194	F—	44		
B-2195	F-			
B-2196	F	HN		
B-2197	F—	HN C		

Examples B-2198 through B-2221 re prepared from Scaffold C-22

	Examples B 2.10	6 tillough B-2221 i	* P	
Example#	R²	Ь,		
B-2198	F—	3.1		
B-2199	F—	S. F.		
B-2200	F—			
B-2201	F—			
B-2202	F-	Z.L		
B-2203	F{}	2		
B-2204	F-_\{	O BR		

Example#	Ŕ²	R ^J		
		0		
B-2205	F—	, O		
B-2206	F—	2,00		
B-2207	F—	3,4		
B-2208	F—		-	
B-2209	F—			
B-2210	F-			
B-2211	F-	F. T.		
B-2212	F—			
B-2213	F—			
B-2214	F-	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		

Example#	Ħ²	R ¹		
B-2215	F—	2000		
B-2216	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
B-2217	F——	LT NH		
B-2218	F-	44		
B-2219	F—			
B-2220	F—	HN		
B-2221	F——}	H. N.		

Example#

B-2226

B-2227

B-2228

860

R2

Examples B-2222 through B-2245 are prepared from Scaffold C-29

R1

Example#

R²

RJ

	· · · · · · · · · · · · · · · · · · ·			
B-2229	s →			
B-2230	s T	22		
B-2231	s T	المراقع المراق		
B-2232	s T			
B-2233	s T			
B-2234	s T			
B-2235	s >	و بره	•	
B-2236	S	N. S.		
B-2237				

Example#

R²

RJ

	·			
B-2238	s →	54 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0		
B-2239		700		
B-2240	S	F 0		
B-2241	S	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		
B-2242	s	Y		
B-2243	s >			
B-2244		HN-O		
B-2245	S	PN O		

Examples B-2246 through B-2269 are prepared from Scaffold C-35

	Examples D-ZZ	46 through B-2269 a	ne prepared	i irom Scar	1010 C-35
Example#		Кı			
B-2246	F-	3,4			
B-2247	F—	ا ا			
B-2248	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\			
B-2249	F—				
B-2250	F—	3,4			
B-2251	F—	2,1			
B-2252	F—	Ş. □ BR			

Example#	R²	H,		
B-2253	F—	بالرا		
B-2254	F—	27		
B-2255	F—	3,4		
B-2256	F—			
B-2257	F——}			
B-2258	F—	7		
B-2259	F—	F 77 0		
B-2260	F—	240		
B-2261	F—			
B-2262	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\		

		· · · · · · · · · · · · · · · · · · ·	 	
Example#	R²	R ^J		
B-2263	F—	10 10 10 10 10 10 10 10 10 10 10 10 10 1		
B-2264	F——}	7, 80 F 0		
B-2265	F—————————————————————————————————————	Y NH		
B-2266	F—	Y		
B-2267	F—			
B-2268	F—	HN-O		
B-2269	F—	N. V.		

866

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Examples B-2270 through B-2317

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In a parallel array reaction block containing 48 fritted vessels, each reaction vessel was charged with 250 mg of polymer bound carbodiimide B48 (1.0 mmol/g resin) and a solution of the acid-containing scaffold c-49 in dimethylformamide (0.1 M, 500 uL). To each slurry was added a solution of pyridine in dichloromethane (0.2 M, 1000 uL) followed by a solution of a unique amine B47 (0.2 M, 375 uL) in dimethylformamide. The reaction mixtures were agitated on a Labline benchtop orbital shaker at 250 RPM for 16-20 h at ambient temperature. The reaction mixtures were filtered into conical vials and the polymer was washed with 1.5 mLdimethylformamide and 2.0 mL of dichloromethane. The filtrates were evaporated to dryness in a apparatus and dimethylformamide (350 uL) was added to each conical vial to dissolve the residue. A solution of tetrafluorophthalic anhydride (1.0 M, 150 uL) in

867

dimethylformamide was added to the reconstituted conical vials and the mixture incubated for 2 hours at ambient temperature. Polyamine polymer B33 (4.0 meg N/g resin, 250 mg) and 1.0 mL dichloromethane was then added to the reaction mixture in each conical vial. After agitating the reaction mixtures for 16 h at 250 RPM on an orbital shaker at ambient temperature, the mixtures were filtered through a polypropylene syringe tube fitted with a porous frit. The polymers were washed twice with dimethylformamide (1.0 mL.each) and the filtrates and 10 washings collected in conical vials. The filtrates were evaporated to dryness and weighed to afford the desired amide products B-2270 through B-2317 as oils or solids. The analytical data and yields for the products prepared in this manner are listed below. 15

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B-2270

B-2271

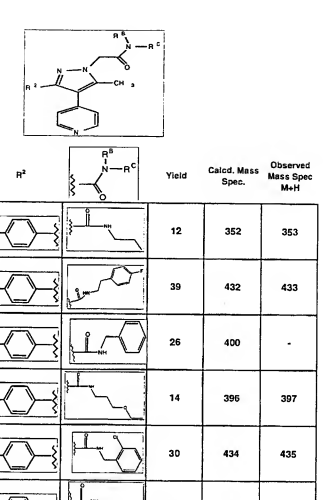
B-2272

B-2273

B-2274

B-2275

B-2276



43

35

443

364

	R ²	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2277		33	490	•
B-2278	F-	53	460	461
B-2279	F-C	10	420	•
B-2280	F—	7	435	436
B-2281		18	401	402
B-2282	F-\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	22	390	413° *M+Na
B-2283	F—N	10	394	417° "M+Na
B-2284		7	423	-
B-2285		23	450	
B-2286		4	506	•

	R²	R ^B N—R ^C	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2287	F—	NH 6	5	437	438
B-2288	F—	i,~~	8	435	436
B-2289	F-		4	450	451
B-2290	F—		9	456	457
B-2291	F—		9	415	416
B-2292	F	NH NH	5	368	369
B-2293	F-\	NH X	5	366	367
B-2294	F-{}) NH	5	381	382
B-2295	F—		16	410	411
B-2296	F-{}		4	483	-

	R²	R ^B N—R ^C	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2297	F—		7	490	
B-2298	F—	المائية المائية	4	537	•
B-2299	F—		4	507	508
B-2300	F—		7	442	-
B-2301	F-		20	396	397
B-2302	F—	اب ا	30	459	•
B-2303	F—{		6	482	
B-2304	F—		5	395	396
B-2305	F-		10	460	•
B-2306	F—	لنب	11	466	467

	R²	RB N—RC	Yleid	Caicd. Mass Spec.	Observed Mass Spec M+H
B-2307	F—		5	421	422
B-2308	F-\{\}		26	470	
B-2309	F-		24	424	425
B-2310	F—		9	348	•
B-2311	F—	o Zi	21	338	339
B-2312	F-{}	NH S	28	398	399
B-2313	F-		6	410	•
B-2314	F—	NH NH	15	363	364
B-2315	F—		11	444	-
B-2316	F—		11	418	•

	R²	HB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2317	F—		36	428	

By analogy to the procedure identified above for the preparation of Examples B-2270 through B-2317, the following examples B-2318 through B-2461 were prepared.

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2318	F—	HN	23	426	427
B-2319	F——}		23	394	•
B-2320	F—		50	490	491
B-2321	F—		49	426	427
B-2322	F—	NH NH	40	366	367
B-2323	F——	0	68	410	411
B-2324	F——}	NH S	57	456	457

	R²		Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2325	F-	NH	41	382	383
B-2326	F—	NH	71	440	441
B-2327	<u>F</u>		36	464	465
B-2328	F—	€ (32	467	468
B-2329	F—	Lo 10	34	465	466
8-2330	F—		26	364	365
B-2331	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	38	464	465
B-2332	F—	N N N	33	483	484
B-2333	F-	NH C	36	378	379

	R²	RB I N RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2334	F-	O NH	44	428	429
B-2335	F	o NH	27	406	407
B-2336	F———	NH NH	41	428	429
B-2337	F—		27	423	424
B-2338	F—		33	469	470
B-2339	F——}	E S	52	518	519
B-2340	F—	NH NH	64	442	443
B-2341	F—	NH	41	350	351
B-2342	F—	NH NH	34	414	415

	T			· · · · · · · · · · · · · · · · · · ·	
	R ²	RB I N-RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2343	F—	O H	29	424	425
B-2344	F-	B r	33	492	493
B-2345	F—	0 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 - 1 -	30	420	421
B-2346	F—	. F	35	474	475
B-2347	F—	DE E	34	392	393
B-2348	F—	NH S	51	458	459
B-2349	F—	0 TO NO.	73	517	518
B-2350	F—		22	448	449
B-2351	F-	, NA	64	486	487

	R²	RB I N-RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2352	<u>F</u>	NH O	41	482	483
B-2353	F——	Zim.	57	438	439
B-2354	F———	**************************************	63	484	485
B-2355	F—		28	536	537
B-2356	F—	, ', ' -Z- -Z-	29	408	409
B-2357	F-\	0 - X - X	41	436	437
B-2358	F——	× × ×	41	451	452
B-2359	F——}	NH O	57	502	503
B-2360	F—	NH NH O	46	496	497

	R²	R ^B I N − R ^c	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2361	F—		13	476	477
B-2362	F—	0 2	46	493	494
B-2363	F—	0=√,	57	396	397
B-2364	F—	0= \	61	438	439
B-2365	F—	0=\\ \-\=\\ \-\=\\ \-\=\\ \-\=\\ \-\=\\ \-\=\\ \\ \-\=\\ \\ \-\=\\ \\ \-\=\\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\ \\	72	424	425

	R²	RB N—R°	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2366	F—		34	380	381
B-2367	F-	0	52	480	481
B-2368	F—	__\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	35	407	407
B-2369	F-\		31	435	436
B-2370	F-		33	414	415
B-2371	F-	~ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \ \	28	366	367
B-2372	F-\		37	422	423

	R²	RB N—Rc	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2373	F—————————————————————————————————————		50	432	433
B-2374	F—{}		29	382	383
B-2375	F—		35	395	396
B-2376	F———		36	428	429
B-2377	F—		68	438	439
B-2378	F—		55	446	447
B-2379	F—	0=\\ \{\frac{1}{2}}	33	364	365
B-2380	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	51	421	422
B-2381	F—		52	429	430

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2382	F C		48	407	408
B-2383			53	382	383
B-2384	F—		38	447	448
B-2385	F—		59	498	450
B-2386	<u>-</u> F—	,	45	429	430
B-2387	F—		74	558	•
B-2388	F—	_N	53	475	-
B-2389	F—	1, N,	33	493	494
B-2390	F		53	487	488

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2391	F—		30	435	436
B-2392	F—		57	464	465
B-2393	F—	~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~ ~	50	418	419
B-2394	F—————————————————————————————————————		65	488	489
B-2395	F—) / S	59	437	438
B-2396	F—	J _N OMe	34	534	535
B-2397	F—	2 N C	32	516	517
B-2398	F——}	N CI	81	533	534
B-2399	F—		55	502	-

	R²	R ^B I N—R ^c	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2400	F—————————————————————————————————————	NH.	34	381	382
B-2401	F—————————————————————————————————————		32	378	379
B-2402	F-\		71	519	520
B-2403	F—S	*	68	527	528
B-2404	F—\}		62	447	448
B-2405	F—————————————————————————————————————	0 / 0	71	536	537
B-2406	F——}		47	394	395
B-2407	F—	~~~~~	65	508	509
B-2408	F—	SME OME	34	495	496

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2409	F-	S S	47	448	449
B-2410	F-__\{		73	542	543
B-2411	F-		81	489	490
B-2412	F—	2-0	54	409	410
B-2413	F—	in the state of th	37	493	494

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec · M+H
B-2414	F-	HN	14	473	474
B-2415	F-	2 Z Z	19	421	422
B-2416	F—	- Z	13	386	387
B-2417	F—		29	414	415
B-2418	F—	D	6	420	421
B-2419	F—————————————————————————————————————	NH CF 3	10	454	-
B-2420	F—	NH NH	5	442	443

	R²	RB N—RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2421	F-_}	CI NH CI	28	454	455
B-2422	F——}	NH O	47	420	421
B-2423	F	NH NH	53	400	401
B-2424	F-		15	400	401
B-2425	F———	F ₃ C	18	522	523
B-2426	F—	B. Z. ←	38	464	465
B-2427	F—————————————————————————————————————		26	468	469
B-2428	F——}	NH S	22	432	433
B-2429	F—	O NH	41	404	405

	R²	R B C C C C C C C C C C C C C C C C C C	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2430	F—	O 2N NO 2	15	476	477
B-2431	F—S	- X	6	446	447
B-2432	F—————————————————————————————————————		37	404	405
B-2433	F—————————————————————————————————————	≥ - ()	8	428	429
B-2434	F—		13	476	477
B-2435	F—		23	442	443
B-2436	F—	 	5	486	487
B-2437	F—		4	492	493
B-2438	F—	\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\\	58	422	423

	R²	RB I N RC	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2439	F———	₹	12	454	455
B-2440	F—	N - N - N - N - N - N - N - N - N - N -	8	521	522
B-2441	F—		6	443	444
B-2442	F—{}		37	514	515
B-2443	F—	, MH	15	518	<u>-</u>
B-2444	F—	J	52	520	-
B-2445	F—	نارز	33	517	518
B-2446	F-	0=30	70	500	501
B-2447	F—	7.5	56	488	489

	R²	RB RC C N C C C C C C C C C C C C C C C C	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2448	F—		51	522	523
B-2449	F—	of E O	19	512	513
B-2450	F———		16	538	539
B-2451	F—		71	511	512
B-2452	F—	TY DOSY	71	500	501
B-2453	F—	2	61	470	•
B-2454	F—		15	472	473
B-2455		N	39	520	•
B-2456	F-		51	533	534

	R²	R ^B − R ^C − R	Yield	Calcd. Mass Spec.	Observed Mass Spec M+H
B-2457	F-	\$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$ \$	55	540	•
B-2458	F—		22	488	489
B-2459	F—	0-√ 0-√ 0-0-€	8	486	487
B-2460	F-	LA CONTRACTOR OF THE CONTRACTO	13	534	53 5
B-2461	F———		13	542	•

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Example C-1

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5-AMINOMETHYL-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

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1-(4-fluorophenyl)-2-(4-pyridyl)-1-ethanone. 4-picoline (40 g, 0.43 mol) was added to a LiHMDS solution (0.45 mol, 450 mL of a 1.0 M solution in THF) over 30 minutes at room temperature (a slight exotherm was observed) The resulting solution was stirred for 1 h. This solution was added to ethyl 4-fluorobenzoate (75.8 g, 0.45 mol, neat) over 1 h. The mixture was stirred overnight (16 h). Water (200 mL) was added and the mixture was extracted with EtOAc (2x200 mL). The organic layer was washed with brine (1x200 mL) and dried over

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Na₂SO₄. The organic layer was filtered and the solvent was removed to leave oily solid. Hexane was added to the oil and the resulting solid was filtered and washed with hexane (cold). A yellow solid was isolated (50 g, 54%):

¹H NMR (CDCl₃) δ 8.58 (d, J = 5.7 Hz, 2H), 8.02 (dd, J = 5.5, 8.0, 2H), 7.12-7.21 (m, 4H), 4.23 (s, 2H);

¹⁹F NMR (CDCl₃) δ -104.38 (m); LC/MS, t_r = 2.14 minutes (5 to 95% acetonitrile/water over 15 minutes at 1 mL/min, at 254 nm at 50°C), M+H = 216; High Resolution MS Calcd for C₂₃H₂₀N₄O₂F (M+H): 216.0825. Found: 216.0830 (Δ mmu = 0.5).

N-benzyloxycarbonyl-5-aminomethyl-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole. A 3L round bottom flask fitted with a mechanical stirrer, No inlet and an addition funnel was was charged with 557 mL (0.56 mol) of 1 M t-BuOK in THF and 53 mL (0.56 mol) of t-BuOH. The ketone, 1 (60 g, 0.28 mol) was dissolved in 600 mL of THF and added to the stirred mixture at room temperature. precipitate formed and the mixture was stirred for 1 h. N-benzyloxycarbonyl-glycinyl N-hydroxysuccinimide (128.6 g, 0.42 mol) was dissolved in 600 mL of THF and added dropwise at r.t. over lh. The mixture was stirred for another 5 minutes and 150 mL of water was added. the pH was adjusted to 6.7 with 70 mL of AcOH. Hydrazine monohydrate (41 mL in100 mL of water) was added via an addition funnel. The mixture was stirred for 1 h and was diluted with 500 mL of water and 500 mL of ethyl acetate. The biphasic mixture was transferred to a sep funnel and the layers were separated. The aqueous layer was extracted with EtOAc (3x300 mL). The organic layer was

WO 00/31063 PCT/US99/26007

dried (Na₂SO₄), filtered and evaporated to leave 157 g of a crude reddish oil.

The oil was suspended in CH2Cl2 and filtered to remove any insoluble material (DCU, hydrazone of the The solution was split into two portions monoketone). and each portion was chromatographed (Biotage 75L, 3% EtOH/CH2Cl2 then 6% EtOH/CH2Cl2). The appropriate fractions were concentrated (some contamination from the monoketone and the hydrazone) from each portion to leave a yellow solid. The solid was suspended in ethyl acetate and heated to boiling for 10 minutes. The solution was allowed to cool to R.T. overnight. The precipitate was filtered to give 30 g of a white solid (27% yield of 2): ¹H NMR (DMF-d₇) δ 13.36 (s, 1H), 8.57 (d, J = 5.8 Hz, 2H), 7.16-7.52 (m, llH), 5.11 (s, 2H), 4.48 (d, J = 5.4 Hz, 2H); ¹⁹F NMR (DMF- d_7) δ -114.9 (m), -116.8 (m) (split fluorine signal is due to the pyrazole tautomers); LC/MS, tr = 3.52 minutes (5 to 95% acetonitrile/water over 15 minutes at 1 mL/min, at 254 nm at 50°C), M+H = 403; High Resolution MS Calcd for $C_{23}H_{20}N_4O_2F$ (M+H): 403.1570. Found: $403.1581 (\Delta mmu = 1.1)$.

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5-aminomethyl-4-(4-pyridyl)-3-(4-fluorophenyl)

pyrazole. To a 1L Parr bottle was added 7 g (17.4 mmol) of 2 and 180 mL of MeOH and 90 mL of THF to give a clear solution. The bottle was purged with nitrogen and 1.5 g of 10% Pd/C (wet Degussa type E101) was added. The Parr bottle was pressured to 40 psi (H₂) and was agitated. Hydrogen uptake was 5 psi after 5 h. The bottle was repressured to 42 psi and was agitated overnight. The bottle was purged with N2 and was filtered through Celite. The Celite was washed with MeOH (3x50 mL) and

the filtrate was concentrated to give 4.5 g of an off-white solid (94%). 1 H NMR (DMSO-d₆) δ 8.52 (d, J = 4.63 Hz, 2H), 7.36 (dd, J = 5.64, 8.1 Hz, 2H), 7.16-7.30 (m, 4H), 3.79 (s, 2H); 19 F NMR (DMSO-d₆) δ -114.56 (m); LC/MS, t_r = 1.21 minutes (5 to 95% acetonitrile/water over 15 minutes at 1 mL/min, at 254 nm at 50°C), M+H = 269 m/z; High Resolution MS Calcd for $C_{15}H_{14}N_4$ F (M+H): 269.1202. Found: 269.1229 (Δ mmu = 2.7).

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The following pyridylpyrazoles (C-2 through C-21, Table C-1) were prepared according to the experimental procedure described above for example C-1.

15

Table C-1.

Exampl	Structure	MW, M +	'H NMR (solvent), ppm
e No.		н	
		Calculat	_
		ed	
		Found	
C-2	N-NH	323.1672	$(DMF-d_7): 8.77 (t, J =$
	F	323.1670	4.4 Hz, 2H), 7.60 (m, 2H),
			7-44 (t, J = 4.4 Hz, 2H),
			7.35 (m, 2H), 3.22 (bd,
			2H), 3.01 (septet, J = 5.3
			Hz, 1H), 2.74 (m, 2H),
			1.95 (m, 4H)

C-4 N-NH NH N	C-3	N-NH	282.127	(DMF-d ₇): 8.77 (br s,
C-4 N-NH NH N	1	F NH ₂	(M)	2H), 7.64-7.62 (m, 2H),
C-4 N-NH NH ₂ 282.127 (DMF-d ₇): 8.77 (br s, 2H), 7.64-7.62 (m, 2H), 7.50 (br s, 2H), 7.38-7.35 (m, 2H), 1.57 (br s, 3H) C-5			282.1245	7.50 (br s, 2H), 7.38-7.34
C-4 N-NH NH ₂ 282.127 (DMF-d ₇): 8.77 (br s, 2H), 7.64-7.62 (m, 2H), 7.50 (br s, 2H), 7.38-7.35 (m, 2H), 4.40-4.37 (m, 1H), 1.57 (br s, 3H) C-5		N.	(M, EI)	(m, 2H), 4.40-4.37 (m,
C-5 N-NH NH2 NN NH2 NN NH2 NN NH2 NN NH3 NH3 NH4	,			1H), 1.56 (br s, 3H)
C-5 N-NH NH2 NN NH2 NN NH2 NN NH4 NN	C-4		282.127	(DMF-d ₇): 8.77 (br s,
C-5 N-NH NH S1 282.1147 (M, EI) (M, EI) 323.1672 (DMSO-d ₆): 8.56 (br, 2H), 7.18 (m, 4H), 2.91 (m, 2H), 2.71 (m, 2H), 1.40 (m, 2H) (m, 2H), 1.40 (m, 2H) (m, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH ₂ NH ₂ N-NH N-NH NH ₂ N-NH NH NH ₂ N-NH NH NH ₂ N-NH NH N			(M)	2H), 7.64-7.62 (m, 2H),
C-5 N-NH 323.1672 323.1687 T.32 (m, 2H), 7.18 (m, 4H), 2.91 (m, 2H), 2.71 (m, 2H), 1.40 (m, 2H) C-6 N-NH 359 (DMSO-d ₆): 8.46 (d, J = 4.6 Hz, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH ₂ 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 2.94 (m, 2H)			282.1147	7.50 (br s, 2H), 7.38-7.35
C-5 N-NH		N I	(M, EI)	(m, 2H), 4.40-4.37 (m,
The state of the s				1H), 1.57 (br s, 3H)
7.32 (m, 2H), 7.18 (m, 4H), 2.91 (m, 2H), 2.71 (m, 2H) 1.88 (m, 1H), 1.65 (m, 2H), 1.40 (m, 2H) C-6 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 4.6 Hz, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH2 313.1465 (DMSO-d ₆): 13.83 (bs, 1.94), 8.61 (d, J = 5.7 Hz, 1.98-1.94)	C-5	^ " \ ^	323.1672	(DMSO-d ₆): 8.56 (br, 2H),
(m, 2H) 1.88 (m, 1H), 1.65 (m, 2H), 1.40 (m, 2H) C-6 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 4.6 Hz, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH2 313.1465 (DMSO-d ₆): 13.83 (bs, 1.94), 8.61 (d, J = 5.7 Hz, 1.98-1.94)		FUTTON	323.1687	7.32 (m, 2H), 7.18 (m,
(m, 2H), 1.40 (m, 2H) C-6 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 4.6 Hz, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH2 313.1465 (DMSO-d ₆): 13.83 (bs, 1.492 (1H), 8.61 (d, J = 5.7 Hz, 1.492 (1H), 9.492 (1H), 9.492 (1H), 9.492 (1H), 9.492 (1H),				4H), 2.91 (m, 2H), 2.71
C-6 N-NH		,,		(m, 2H) 1.88 (m, 1H), 1.65
359 4.6 Hz, 2H), 7.32-7.13 (m, 7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH2 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98- 2.94 (m, 2H) C-8 N-NH NH2 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 (1H), 8.61 (d, J = 5.7 Hz, 313.1492 (1H), 8.61 (d, J = 5.7 Hz,		•		
7H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH ₂ 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 (1H), 8.61 (d, J = 5.7 Hz, 414.1492 (1H), 8.61 (d, J = 5.7 Hz, 414	C-6		359	$(DMSO-d_6): 8.46 (d, J =$
4.06 (t, J = 7.0 Hz, 1H), 2.98-2.95 (m, 2H) C-7 N-NH NH ₂ 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98- 2.94 (m, 2H) C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 1H), 8.61 (d, J = 5.7 Hz,			359	
2.98-2.95 (m, 2H) C-7 N-NH NH ₂ 359 (DMSO-d ₆): 8.46 (d, J = 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 1H), 8.61 (d, J = 5.7 Hz, 413.1492 1H)	ļ			7H), 6.98-6.96 (m, 4H),
C-7 N-NH				4.06 (t, J = 7.0 Hz, 1H),
359 5.4 Hz, 2H), 7.32-7.28 (m, 2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98-2.94 (m, 2H) C-8 N-NH NH _Z 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 1H), 8.61 (d, J = 5.7 Hz, 313.1492 1H)				2.98-2.95 (m, 2H)
2H), 7.20-7.12 (m, 5H), 6.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98- 2.94 (m, 2H) C-8 N-NH 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 lH), 8.61 (d, J = 5.7 Hz,	C-7	N-NH NH ₂	359	$(DMSO-d_6): 8.46 (d, J =$
C-8 N-NH 0.98-6.96 (m, 4H), 4.06 (t, J = 7.0 Hz, 1H), 2.98- 2.94 (m, 2H) C-8 N-NH NHz 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 1H), 8.61 (d, J = 5.7 Hz,		F	359	5.4 Hz, 2H), 7.32-7.28 (m,
(t, $J = 7.0 \text{ Hz}$, IH), $2.98-2.94 \text{ (m, } 2H)$ C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 IH), 8.61 (d, $J = 5.7 \text{ Hz}$,				2H), 7.20-7.12 (m, 5H),
2.94 (m, 2H) C-8 N-NH 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 1H), 8.61 (d, J = 5.7 Hz,		1		6.98-6.96 (m, 4H), 4.06
C-8 N-NH NH ₂ 313.1465 (DMSO-d ₆): 13.83 (bs, 313.1492 lH), 8.61 (d, J = 5.7 Hz,				(t, J = 7.0 Hz, 1H), 2.98-
NH_2 313.1492 1H), 8.61 (d, J = 5.7 Hz,				_
	C-8		313.1465	
OCH,		F COCH	313.1492	1H), 8.61 (d, $J = 5.7$ Hz,
2H), 8.33 (bs, 1H), 7.33	1	N N		
(m, 6H), 4.44 (m, 1H),				
3.63 (m, 2H), 3.27 (s, 3H)				3.63 (m, 2H), 3.27 (s, 3H)

C-10 Note	C-9	N-NH N-NH	313.1465	$(DMSO-d_6): 8.55 (dd, J =$
7.32 (m, 2H), 7.26 (dd, J = 1.6, 4.4 Hz, 2H), 7.22-7.16 (m, 2H), 4.06 (t, J = 6.5 Hz, 1H), 3.49 (d, J = 6.6 Hz, 2H), 3.20 (s, 3H) C-10 N-NH NH, 354 CONHICK 1354 CONHICK 1355 CONHICK 1355 CONHICK 1356 CONHICK 1357 CONHICK 1358 CONHICK 1458 CONHICK 1558 CONHICK 1658 CONHI			313.1457	1.5, 4.4 Hz, 2H), 7.37-
7.16 (m, 2H), 4.06 (t, J = 6.5 Hz, 1H), 3.49 (d, J = 6.6 Hz, 2H), 3.20 (s, 3H) C-10				7.32 (m, 2H), 7.26 (dd, J
C-10		,		= 1.6, 4.4 Hz, 2H), 7.22-
C-10 N-NH NH2 STATE C-10 N-NH NH2 STATE CONHICH CONH				7.16 (m, 2H), 4.06 (t, J =
C-10 C-10 CONHICK 354 (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (dt, J=7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-11 C-11 CONHICK 354 (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 C-13 C-13 C-14 C-15				6.5 Hz, 1H), 3.49 (d, J =
This is the content of the content o			-	6.6 Hz, 2H), 3.20 (s, 3H)
CONHCH Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-11 N-NH NH SO (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH 283.1359 (DMSO-d ₆): 8.53 (d, J= 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83 (d, J= 6.0 Hz, 2H) C-13 NANH NH 297.1515 (DMSO-d ₆): 8.53 (d, J= 5.4 Hz, 2H), 7.34 (dd, J= 5.4 Hz, 2H), 7.34 (dd, J= 5.4 Hz, 2H), 7.34 (dd, J=	C-10	N-NH NH ₂	354	(DMSO-d ₆): 13.03 (bs,
Hz, 2H), 7.58 (Bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH		FUIT	354	1H), 8.50 (dd, J=1.6, 2.7
T.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-11 N-NH CONHCH, 354 (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH 283.1359 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83 (d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J =		CN CONHCH		Hz, 2H), 7.58 (bq, J=4.3
(t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-11 N-NH NH 354 (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH 283.1359 DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H), 7.34 (dd, J =				Hz, 1H), 7.3 (m, 2H),
(d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-11 N-NH NH CONHICH STATE (DMSO-d6): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH 283.1359 (DMSO-d6): 8.53 (d, J= 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J=6.0 Hz, 2H), 2.77 (d, J=6.0 Hz, 2H) C-13 N-NH NH 297.1515 (DMSO-d6): 8.53 (d, J= 5.4 Hz, 2H), 7.34 (dd, J=				7.12-7.21 (m, 4H), 3.77
C-11 N-NH NH ₂ CONHCH ₃ CONHCH ₄ N-NH NH ₂ CONHCH ₃ (dt, J=7.3, 7.1 Hz, 2H) (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH ₂ 283.1363 C-13 N-NH NH ₂ 283.1363 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.21-7.17 (m, 4H), 2.83 (d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH NH ₂ 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2				(t, J= 6.3 Hz, 1H), 2.45
C-11 N-NH NH CONHICH3 (dt, J=7.3, 7.1 Hz, 2H) (DMSO-d ₆): 13.03 (bs, 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH 283.1359 283.1363 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J =				(d, J=4.5 Hz, 3H), 1.97
C-11 N-NH NH CONHCH; 354 354 1H), 8.50 (dd, J=1.6, 2.7 Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH 283.1359 283.1363 CDMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H) C-13 N-AH NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H), 7.34 (dd, J =				(t, J= 7.4 Hz, 2H), 1.85
This is the second of the seco				(dt, J=7.3, 7.1 Hz, 2H)
Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J=6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J=7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH NH _z 283.1359 283.1363 C-12 N-NH NH _z 283.1363 C-13 N-NH NH _z 283.1359 283.1363 C-14 C-15 N-NH NH N	C-11	N-NH NH ₂	354	(DMSO-d ₆): 13.03 (bs,
Hz, 2H), 7.58 (bq, J=4.3 Hz, 1H), 7.3 (m, 2H), 7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH 283.1359 283.1363 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H)			354	1H), 8.50 (dd, J=1.6, 2.7
7.12-7.21 (m, 4H), 3.77 (t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH 283.1359 283.1363 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 27.77 (d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H)		N CONNERS		Hz, 2H), 7.58 (bq, J=4.3
(t, J= 6.3 Hz, 1H), 2.45 (d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) (DMSO-d ₆): 8.53 (d, J = 283.1363				Hz, 1H), 7.3 (m, 2H),
(d, J=4.5 Hz, 3H), 1.97 (t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 NHz 283.1359 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-AH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H)				7.12-7.21 (m, 4H), 3.77
(t, J= 7.4 Hz, 2H), 1.85 (dt, J=7.3, 7.1 Hz, 2H) C-12 N-NH 283.1359 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83 (d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz,				(t, J= 6.3 Hz, 1H), 2.45
(dt, J=7.3, 7.1 Hz, 2H) C-12 F NH ₂ 283.1359 (DMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 6.0 Hz, 2H) C-13 S-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H)				(d, J=4.5 Hz, 3H), 1.97
C-12 NH _Z 283.1359 283.1363 CDMSO-d ₆): 8.53 (d, J = 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83 (d, J = 6.0 Hz, 2H) C-13 N-AH NH _Z 297.1515 (DMSO-d ₆): 8.53 (d, J = 6.0 Hz, 2H) C-13 SAME 297.1515 (DMSO-d ₆): 8.53 (d, J = 5.4 Hz, 2H), 7.34 (dd, J = 5.4 Hz, 2H), 7.34 (dd, J = 6.0 Hz, 2H)	1			(t, J= 7.4 Hz, 2H), 1.85
283.1363 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH	1			
283.1363 5.0 Hz, 2H), 7.37-7.32 (m, 2H), 7.21-7.17 (m, 4H), 2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 297.1515 5.4 Hz, 2H), 7.34 (dd, J =	C-12	~ // \	283.1359	· ·
2.83(d, J = 6.0 Hz, 2H), 2.77 (d, J = 6.0 Hz, 2H) C-13 N-AH 297.1515 (DMSO-d ₆): 8.53 (d, J = 297.1515 5.4 Hz, 2H), 7.34 (dd, J =			283.1363	5.0 Hz, 2H), 7.37-7.32 (m,
2.77 (d, J = 6.0 Hz, 2H) C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 297.1515 5.4 Hz, 2H), 7.34 (dd, J =		N.		2H), 7.21-7.17 (m, 4H),
C-13 N-NH 297.1515 (DMSO-d ₆): 8.53 (d, J = 297.1515 5.4 Hz, 2H), 7.34 (dd, J =				1
297.1515 5.4 Hz, 2H), 7.34 (dd, J =				I
	C-13	N-NH NH2	Í	
5.8, 8.2 Hz, 2H), 7.18			297.1515	
		N. P		5.8, 8.2 Hz, 2H), 7.18

			(dd, J = 5.8, 9.8 Hz, 4H),
			2.68 (t, $J = 7.3 \text{ Hz}, 2H$),
			2.52 (m, 2H), 1.64 (m, 2H)
C-14	CI N-NH NH2	284.0829	(CD ₃ OD): 8.74 (br, 2H),
	T NITZ	284.0806	7.77 (br, 2H), 7.45-7.58
			(m, 3H), 7.30-7.40 (m,
	N.		1H), 4.43 (s, 2H)
C-15	N-NH NH2	285	(DMSO-d ₆): 8.53 (br, 2H),
	a-1	285	7.56 (br, 2H), 7.26 (m,
			4H), 3.75 (br, 2H)
C-16	N-NH N-NH	329, 331	$(DMSO-d_6): 8.53 (d, J =$
	Br	329, 331	4.4 Hz, 2H), 7.42 (d, J =
			7.9 Hz, 2H), 7.34 (d, J = 1)
	N		8.5 Hz, 2H), 7.24 (d, J =
			4.6 Hz, 2H), 3.76 (bs, 2H)
C-17	CI N-NH	339	$(DMSO-d_6): 8.53 (t, J =$
	NH	339	4.3 Hz, 2H), 7.33 (m, 3H),
			7.19.(t, J = 4.6 Hz, 2H),
	N		7.14 (d, J = 7.3 Hz, 1H),
	Ì		3.23 (m, 2H), 2.88, (m,
			3H), 1.92, (m, 3H), 1.70
			(m, 1H)
C-18	N-NH	339	$(DMSO-d_6): 8.57 (d, J =$
1	CI NH	339	4.6 Hz, 2H), 7.41 (d, J =
			8.3 Hz, 2H), 7.29 (d, J =
			8.5 Hz, 2H), 7.20 (d, J =
			4.8 Hz, 2H), 3.18 (bd,
			2H), 2.88 (m, 1H), 2.76
			(m, 2H), 1.82 (br, 4H)
C-19	N-NH	383, 385	(DMSO-d ₆): 8.56 (br, 2H),
	Br	383, 385	7.52 (br, 2H), 7.14-7.29
		·	(m, 4H), 2.99 (br, 2H),

2.71 (br, 1H), 2.51 (br,
2H), 1.68 (br, 4H)

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The following pyridylpyrazoles (C-22 through C-40, Table C-2) are prepared utilizing the general schemes C-1 and C-2 and the experimental procedure described for example 15. C-1 above.

Table C-2

Cmpd. No.	Structure
C-22	N-NH NHR S
C-23	F-NH NH
C-24	F NH

C-25	Br N-NH NH2
C-26	H ₃ C N-Ni+ NH ₂
C-27	Br. N-NH
C-28	H ₂ C N-NH
C-29	N-NH NH ₂
C~30	N-NH NN
C-31	F ₃ C N-NH
C-32	F-NH NH2
C-33	F-V-NH

C-34	F-NH NH2
C-35	F-O-NH
C-36	F-O-NH2
C-37	F-NH2
C-38	F-V-NH
C-39	F-O-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-N-
C-40	F-NH CO+Bu
C-41	P H NH
C-42	N-NH H NH
C-43	F HN
C-44	P-NH H

C-45	F-NH H
C-46	F CH,
C-47	F CH3
C-48	F-WH-CH,

Example C-49

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Step A

The pyrazole (2.60 g, 10.3 mmol) from example $_{\rm C-4}$ was suspended in 52 mL of dichloroethane and 52 mL of 2.5 M

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Tetrabutylammonium hydroxide (0.5 mL of a 1 M acueous solution) was added to the stirred mixture. this mixture was added t-butyl bromoacetate (2.10 g, 10.8 The reaction mixture was stirred at room temperature for 4 h. The mixture was poured onto 200 mL of CH2Cl2 and 200 mL of H2O. The phases were separated and the organic phase was washed with water (1x100 mL) and brine (1x100 mL). The organic layer was dried over Na2SO4 and was filtered. The solvent was removed to leave an off-white solid. This solid was triturated with hexane and the resulting solid isolated by filtration. The solid was washed with hexane to leave 3.4 g of a white solid (90%).

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Step B

The alkylated pyrazole (3.7 g, 10.1 mmol) from Step A was treated with 57 mL of 4 N HCL in dioxane. The solution was stirred at room temperature for 4 h. The solvent was removed under reduced pressure and residue was dissolved in THF. The solution was treated with propylene oxide (10.3 mmol) and was stirred for 1h at room temperature. The solvent was removed to leave an The residual solvent was chased with several portions of EtOH. The resulting solid was triturated Example C-49 was with Et₂O and the title compound isolated by filtration to afford 3.0 g of an off-white 30 solid (95%). Mass spec: M+H cald: 312; found 312. NMR (DMSO-d6): 8.81 (d, J = 6.4 Hz, 2H), 7.73 (d, J =

5.8 Hz, 2H), 7.40 (m, 2H), 7.23 (t, J = 8.5 Hz, 1H), 5.16 (s, 2H), 2.40 (s, 3H).

Example C-50

According to the procedure described above in Example C
49, Example C-50 was also prepared starting from 4-[3-(4-fluorophenyl)-1H-pyrazole-4-yl]pyridine. Mass spec: M+H cald: 298; found 298.

14 NMR (DMSO-d6): 8.75 (d, J = 6.4 Hz, 2H), 8.68 (s, 1H), 7.78 (d, J = 6.6 Hz, 2H), 7.52 (dd, J = 5.4, 8.5 Hz, 2H), 7.31 (t, J = 8.9 Hz, 2H), 5.16 (s, 2H).

Example C-51

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Starting with the N-Boc-piperidinyl analog of Example C-2, Example C-51 is also prepared according to the methods described in Scheme C-1.

Example C-52

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Step A: Picoline is treated with a base chosen from but not limited to n-BuLi, LDA, LiHMDS, tBuOK, or NaH in an organic solvent such as THF, ether, t-BuOH or dioxane from -78 °C to 50 °C for a period of time from 10 minutes to 3 hours. The picoline solution is then added to a solution of N-Cbz-(L)-phenylalaninyl N-hydroxysuccinimide. The reaction is allowed to stir from 30 minutes to 48 hours during which time the temperature may range from -20 °C to 120 °C. The mixture is then poured into water and extracted with an organic solvent. After drying and removal of solvent the pyridyl monoketone is isolated as a crude solid which could be purified by crystallization and/or chromatography.

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25 Step B: A solution of the pyridyl monoketone in ether, THF, tBuOH, or dioxane is added to a base chosen from but

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not limited to n-BuLi, LDA, LiHMDS, tBuOK, or NaH contained in hexane, THF, ether, dioxane, or tBuOH from - 78 °C to 50 °C for a period of time from 10 minutes to 3 hours. Formyl acetic anhydride is then added as a solution in THF, ether, or dioxane to the monoketone anion while the temperature is maintained between -50 °C and 50 °C. The resulting mixture is allowed to stir at the specified temperature for a period of time from 5 minutes to several hours. The resulting pyridyl diketone intermediate is utilized without purification in Step C.

Step C: The solution containing the pyridyl diketone is quenched with water and the pH is adjusted to between 4 and 8 utilizing an inorganic or organic acid chosen from HOAC, H₂SO₄, HCl, or HNO₃. The temperature during this step is maintained between -20 °C and room temperature. Hydrazine or hydrazine hydrate is then added to the mixture while maintaining the temperature between -20 °C and 40 °C for a period of 30 minutes to several hours. The mixture is then poured into water and extracted with an organic solvent. The N-Cbz-protected pyridyl pyrazole is obtained as a crude solid which is purified by chromatography or crystallization.

5 Step: D

The CBZ protecting group is cleaved using hydrogen gas under pressure and Pd-C in an alcohol solvent, affording scaffold C-52 after filtration and concentration.

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The following compounds C-53 through C-59 in Table C-3 are prepared according to the general procedure described above for the preparation of C-52.

Table C-3

Example No.	Structure
C-53	H ₂ N H

C-54	N-NH H ₂ N Boc
C-55	H ₂ N N-NH N Boc
C-56	H ₂ N N-NH H
C-57	H ₂ N N-NH H
C-58	H₂N N-NH NH-Boc
C-59	H ₂ N N-NH NH-Box

Example C-60

5 Step A:

A Boc protected pyridylpyrazole is treated with benzaldehyde in methylene chloride at room temperature in

the presence of a drying agent for a period of time ranging from 1-24 h. Solvent is then evaporated and the resulting imine is used in step B without further purification.

Step B:

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The pyridylpyrazole imine is dissolved in THF and stirred under nitrogen at temperatures ranging from -78 to -20 °C. A base such as LDA, n-BuLi, or LiHMDS is added dropwise to the mixture which is then stirred for an additional 10 minutes to 3 h. Two equivalents of a methyl iodide are then added to the mixture and stirring is continued for several hours. The mixture is then quenched with acid and allowed to warm to room temperature and stirred several hours until cleavage of the Boc and the imine functions is complete. The pH is adjusted to 12 and then the mixture is extracted with an organic solvent, which is dried and evaporated. The crude pyridylpyrazole is then crystallized and/or chromatographed to give purified C-60.

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1) Base 2) Mel 3) Acid, H₂O

C-60

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Example C-61

10 Example C-61 is prepared according to the method described in example C-60, substituting 1,4-dibromobutane for methyl iodide.

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Example C-62 is prepared according to the method described in example C-60, substituting 1,3-dibromoethane for methyl iodide.

Example C-63

The synthesis of compound C-63 starts with the condensation reaction of bromomaleic anhydride B77 with 10 2, 4-dimethoxybenzylamine in acetic acid and acetic anhydride. The maleimide B78 is then treated with 4'fluoroacetophenone in the presence of catalytic amount Pd₂(dba)₃ and sodium t-butoxide to form the 15 fluoroacetophenone substituted maleimide B79. B79 is then treated with tert-butoxybis(dimethylamino)methane to yield the a-ketoenamine B80. The a-ketoenamine B80 is condensed with hydrazine to form the N-protected maleimide pyrazole B81. The 2,4-dimethoxybenzyl group is cleaved with ceric ammonium nitrate (CAN) to give the 20 title compound C-63.

Example C-64

Using the method described in Schemes C-6 and C-7, 10 Example 64 is prepared.

Example C-65

Using the method described in Schemes C-6 and C-7, Example 65 is prepared.

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Example C-66

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Using the method described in Schemes C-6 and C-7, Example C-66 is synthesized, substituting N-2,4-20 dimethoxybenzyl-4-bromopyridone for B78.

Example C-67

F N-NH NH₂

Using the method described in Schemes C-6 and C-7, Example C-67 is synthesized, substituting N-2,4-10 dimethoxybenzyl-4-bromopyridone for B78, and substituting N-Boc-glycyl N-hydroxysuccinimide for B82.

Example C-68

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Using the method described in Schemes C-6 and C-7, 20 Example C-68 is synthesized, substituting N-2,4-dimethoxybenzyl-4-bromopyridone for B78.

Example C-69

Using the method described in Schemes C-6 and C-7, Example 69 is prepared, substituting N-Boc-nipecotyl N-hydroxysuccinimide for B83.

Example C-70

Using the method described in Schemes C-6 and C-7, Example 70 is prepared, substituting N-Boc-nipecotyl Nhydroxysuccinimide for B83.

Example C-71

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Using the method described in Schemes C-6 and C-7, Example 71 is prepared, substituting N-methyl-3-bromomaleimide for B78.

Example C-72

10 Using the method described in Schemes C-6 and C-7, Example 72 is prepared, substituting N-methyl-3-bromomaleimide for B78, and substituting N-Boc-nipecotyl N-hydroxysuccinimide for B83.

Example C-73

Using the method described in Schemes C-6 and C-7,
20 Example 73 is prepared, substituting N-methyl-3-bromomaleimide for B78 and substituting N-Boc-nipecotyl
N-hydroxysuccinimide for B83.

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General Synthetic Procedures

Scheme C-8 illustrates a general method that can be used for the introduction of various groups on an unsubstituted nitrogen atom that is present as part of pyrazole (Cviii) with appropriately substituted aldehydes (R₁₀₂CHO) or ketones (R₁₀₂COR₁₀₃) in the presence of a reducing agent such as sodium cyanoborohydride or sodium triacetoxyborohydride affords the desired products (Cix). Typical conditions for the reductive alkylation include the use of an alcoholic solvent at temperatures ranging from 20 °C to 80 °C. In Scheme C-8, R₁₀₂ and R₁₀₃ are selected from but not limited to alkyl, benzyl, substituted benzyl, arylalkyl, heteroarylalkyl.

Scheme C-9 illustrates another method for introduction of substituents on the unsubstituted nitrogen atom present as part of the C-3 position of the pyrazole (Cviii). Treatment of the pyrazole (Cviii) with

a suitable alkylating agent (R_{10},X) such as an alkyl chloride, alkyl bromide, alkyl iodide or with an alkyl methanesulfonate or alkyl p-toluenesulfonate in the presence of a suitable base affords the desired alkylated pyrazoles (Cx). Examples of suitable bases include diisopropylethylamine, triethylamine, N-methylmorpholine, potassium carbonate and potassium bicarbonate.

Scheme C-9

Typical conditions for the alkylation include reaction with the suitable base in a polar aprotic solvent such as acetonitrile, dimethylformamide, dimethylacetamide or dimethyl sulfoxide at temperatures ranging from 20 °C to 150 °C. Typical R₁₀₄ substituents are selected from but are not limited to alkyl, substituted benzyl, heteroaromatic, substituted heteroalkyl and substituted heteroarylalkyl groups.

Compounds containing acyl, sulfonyl or ureidyl groups at the nitrogen atom can be prepared as shown in Scheme C-10. Treatment of the pyrazole Cviii with a suitable acylating agent in the presence of a base such as N-methylmorpholine, triethylamine, diisopropylethylamine or dimethylamino pyridine in an

organic solvent such as dichloromethane, dichloroethane or dimethylformamide at temperatures ranging from 20 °C to 120 °C affords the desired acylated pyrazoles (Cxi). Suitable acylating agents include acid halides, activated esters of acids such as the N-hydroxysuccinimde esters, p-nitrophenyl esters, pentafluorophenyl esters, sulfonyl halides, isocyanates, and isothiocyanates.

Scheme C-10

A general synthesis of 2-substituted pyrimidinylpyrazole compounds of type Cxv is shown in Scheme C-11.

Step A:

4-Methyl-2-methylmercaptopyrimidine is treated with a base selected from but not limited to n-BuLi, LDA, LiHMDS, t-BuOK, NaH in an organic solvent such as THF, ether, t-BuOH, dioxane from -78 °C to 50 °C for a period of time from 30 minutes to 5 hours. The resulting 4-methyl anion is then added to a solution of an appropriate ester 888. The reaction is allowed to stir from 30 minutes to 48 hours during which time the

temperature may range from 0 °C to 100 °C. The reaction mixture is then poured into water and extracted with an organic solvent. After drying and removal of solvent the desired monoketone B89 is isolated as a crude solid which can be recrystallized or purified by chromatography.

Step B:

Monoketone B89 is treated with a base selected from but not limited to n-BuLi, LDA, LiHMDS, t-BuOK, NaH, K,CO, or Cs,CO, in an organic solvent such as THF, ether, t-BuOH, dioxane, toluene or DMF from -78 °C to 50 °C for a period of time from 30 minutes to 5 hours. A solution of an appropriately activated ester of a carboxylic acid CbzNR*-(CH₂)_nCR*(R°)-COOH or BocNR*-(CH₂)_nCR*(R°)-COOH, preferably but not limited to the N-hydroxysuccinimide ester B90 is then added to the monoketone anion while maintaining the temperature between 0 °C to 100 °C. The reaction is allowed to stir at the specified temperature for a period of time ranging from 30 minutes to 48 hours. The resulting pyrimidine diketone intermediate B91 is utilized without further purification in Step C.

Step C:

The solution or suspension containing the diketone intermediate B91 is quenched with water and the pH adjusted to between 4 and 8 using an acid chosen from AcOH, H,SO₄, HCl or HNO, while maintaining the temperature between 0 °C to 40 °C. Hydrazine or hydrazine monohydrate is then added to the mixture while maintaining the temperature between 0 °C to 40 °C. The mixture is stirred

for a period of 30 minutes to 16 hours maintaining the temperature between 20 °C to 50 °C, poured into water and extracted with an organic solvent. The pyrimidinyl pyrazole CxiiBoc or CxiiCbz is obtained as crude solid which is purified by chromatography or crystallization.

Step D:

the pyrimidinyl 2-methylmercapto group in pyrazole (CxiiBoc or CxiiCbz) is oxidized to the 2methylsulfone (where n = 2) or the 2-methylsulfoxide (where n = 1) using either Oxone or m-chloroperbenzoic acid as an oxidizing agent in a suitable solvent at temperatures ranging from 25 °C to 100 °C. Solvents of oxidation include dichloromethane, for the choice acetonitrile, tetrahydrofuran or hydroalcoholic mixtures. The 2-methylsulfone (n = 2) or the 2-methylsulfoxide (n =1) (CxiiiBoc or CxiiiCbz) is purified by crystallization or chromatography.

Step E:

2-methylsulfone/2-methylsulfoxide group in or CxiiiCBz is conveniently displaced with CxiiiBoc various amines or alkoxides at temperatures ranging from 20 °C to 200 °C in solvents that include but are not limited to dimethylformamide, acetonitrile. tetrahydrofuran and dioxane. The alkoxides can be generated from their alcohols by treatment with a base selected from but not limited to sodium hydride, lithium hexamethyldisilazide, potassium tertiary-butoxide solvents such as tetrahydrofuran, dimethylformamide and

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dioxane at temperatures ranging from 0 °C to 100 °C. The resulting 2-amino or 2-oxo derivatives (CxivBoc or CxivCbz) are purified by either chromatography or crystallization.

Step F:

The carbamate protecting groups from CxivBoc or CxivCbz are removed to afford the desired compounds Cxv containing either a free primary amine $(R^{\kappa}$ is hydrogen) or a free secondary amine (R" is not equal to hydrogen). Boc protecting groups are cleaved utilizing either chloride trifluoroacetic acid in methylene or hydrochloric acid in dioxane at room temperature for several hours. The Cbz protecting groups are cleaved using hydrogen gas at atmospheric or higher pressures and a catalyst (palladium on charcoal) in an alcoholic solvent. The resulting amines Cxv are then crystallized or purified by chromatography.

SCHEME C-11

Cxv

The following examples contain detailed descriptions of the methods of preparation of compounds that form part of the invention. These descriptions are presented for illustrative purposes only and are not intended as a restriction on the scope of the invention. All compounds showed NMR spectra consistant with their assigned structures.

Example C-74

5-(4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

Example C-1 and of following the method By substituting methyl-4-chlorobenzoate for ethyl-4fluorobenzoate and N-t-butoxycarbonyl-isonipecotyl Nhydroxysuccinimide for N-benyloxycarbonyl-glycinyl Nhydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound as the hydrochloride salt: $^{1}HNMR$ (d_e-DMSO) δ 8.57 (d, J = 4.83 Hz, 2 H), 7.41 (d, J = 8.26 Hz, 2 H), 7.29 (d, J =8.26 Hz, 2 H), 7.20 (d, J = 4.63 Hz, 2 H), 3.18 (bd, J =

12.08 Hz, 2 H), 2.88 (m, 1 H), 2.76 (m, 2 H), 1.82 (bs, 4 H). MS (M+H): 339 (base peak).

Example C-75

5-(N-METHYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

To a solution of 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) (25 g, 61 mmol) in 140 mL of formic acid (96%) was added 50 g of formaldehyde (37%). The solution was stirred at 75 °C for 48 h and was cooled to room temperature. The excess formic acid was removed under reduced pressure and the residue was dissolved in 100 mL of water. The solution was added to concentrated NH₄OH/H₂O and the mixture was extracted with ethyl acetate (3 x 200 mL). The combined organic layers were washed with brine (1 x 250 mL) and was dried over Na₂SO₄. The solution was filtered and concentrated to leave a white solid. The solid was triturated with ether and was filtered to afford the title compound: MS (M+H): 353 (base peak).

Example C-76

5-(N-ACETYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

To a stirred suspension of 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) (1 g, 2.4 mmol) in 24 mL of CH₂Cl₂ was added 4-dimethylamino pyridine (0.88 g, 7.2 mmol) and acetyl chloride (0.21 g, 2.6 mmol). The solution was stirred for 3 h and the solvent was removed under reduced pressure. The residue was treated with saturated NH₂OH (20 mL) and the suspension was extracted with ethyl acetate (3 x 30 mL). The combined extracts were washed with brine (1 x 50 mL), dried over MgSO₄, filtered and concentrated to leave a solid. The solid was triturated with ether and was filtered to leave the title compound: MS (M+H): 381 (base peak).

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Example C-77

5-(N-METHOXYACETYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting methoxy acetyl chloride for acetyl chloride the title compound was prepared: 1 HNMR (DMSO- d_{s}) δ 8.75 (d, J = 6.72 Hz, 2 H), 7.70 (d, J = 6.72 Hz, 2 H), 7.38 (d, J = 8.60 Hz, 2 H), 7.29 (dd, J = 6.72, 1.88 Hz, 2 H), 4.40 (d, J = 11.8 Hz, 1 H), 4.05 (m, 2 H), 3.70 (d, J = 12.70 Hz, 1 H), 3.25 (s, 3 H), 3.0 (m, 2 H), 2.55 (m, 1 H), 1.7 (m, 4 H). MS (M+H): 411 (base peak).

Example C-78

5-(N-METHYLSULFONYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting methylsulfonyl chloride (2.0 equivalents) for acetyl chloride the title compound was prepared: 1 HNMR (DMSO- d_{s}) δ 8.70 (d, J = 6.72 Hz, 2 H), 7.72 (d, J = 6.72 Hz, 2 H), 7.38 (d, J = 7.66 Hz, 2 H), 7.30 (dd, J = 6.72, 1.88 Hz, 2 H), 3.58 (bd, J = 11.8 Hz, 2 H), 2.87 (m, 1 H), 2.82 (s, 3 H), 2.72 (m, 2 H), 1.85 (m, 4 H). MS (M+H): 417 (base peak).

Example C-79

5-[N-METHOXYETHYL-4-PIPERIDYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

To a stirred suspension of 5-(4-piperidy1)-4-(4-pyridy1)-3-(4-chloropheny1) pyrazole hydrochloride (Example C-74) (500 mg, 1.2 mmol) in 12 mL of DMF was added Hunig's base (790 mg, 6.1 mmol) and 2-bromoethy1 methy1 ether (850 mg, 6.1 mmol). The solution was stirred at room temperature for 5 days. The solution was poured onto 2.5 N NaOH and was extracted with ethy1 acetate (3 x 100 mL). The combined extracts were washed with water (3 x 100 mL) and brine (1 x 100 mL). The organic phase was dried over Na₂SO₄ and was filtered. The

solvent was removed under reduced pressure to leave a solid. The solid was triturated and filtered to leave the title compound: 1 HNMR (CDCl₁) δ 8.63 (d, J = 4.23 Hz, 2 H), 7.28 (m, 4 H), 7.14 (d, J = 4.43 Hz, 2 H), 3.57 (t, J = 5.24 Hz, 2 H), 3.38 (s, 3 H), 3.14 (bd, J = 10.1 Hz, 2 H), 2.79 (m, 1 H), 2.68 (t, J = 5.04, 2 H), 2.08 (m, 4 H), 1.92 (m, 2 H). MS (M+H): 397 (base peak).

Example C-80

5-(N-ALLYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of example C-79 and substituting allyl bromide for 2-bromoethyl methyl ether the title compound was prepared: MS (M+H): 379 (base peak)

Example C-81

5-(N-PROPARGYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of example C-79 and substituting propargyl bromide for 2-bromoethyl methyl ether the title compound was prepared: MS (M+H): 377 (base peak)

Example C-82

5-[N-(2-METHYLTHIAZOLYL)-4-PIPERIDYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

To a suspension of 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) in 12 mL of MeOH was added trimethyl orthoformate (2.6 g.

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24.4 mmol) and 2-thiazolecarboxaldehyde (1.4 g, 12.2 mmol). The suspension was stirred at room temperature for 2 h. To this mixture was added NaCNBH, (1.5 g, 24.4 mmol) and the resulting suspension was stirred at room temperature for 7 days. The mixture was poured onto 2.5 N NaOH and was extracted with ethyl acetate (2 x 100 mL). The combined extracts were washed with brine (1 x 100 mL), dried over Na,SO4, filtered and concentrated to leave a solid. This solid was triturated with ether and filtered to afford the title compound: MS (M+H): 436 (base peak).

Example C-83

5-(4-PIPERIDYL)-4-(4-PYRIDYL)-3-[4-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-(trifluoromethyl)benzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonyl-isonipecotyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate

was accomplished with 4 N HCl in dioxane to afford the title compound as its hydrochloride salt: MS (M+H): 373 (base peak).

Example C-84

5-(N-METHYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-[4-(TRIFLUOROMETHYL) PHENYL] PYRAZOLE

By following the method of Example C-75 and substituting 5-(4-piperidy1)-4-(4-pyridy1)-3-[4-(trifluoromethyl)phenyl] pyrazole hydrochloride (Example C-83) for 5-(4-piperidy1)-4-(4-pyridy1)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 387 (base peak).

Example C-85

5-[N-(2-PROPYL)-4-PIPERIDYL]-4-(4-PYRIDYL)-3-[4-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

To a solution of 5-(4-piperidyl)-4-(4-pyridyl)-3-[4-(trifluoromethyl)phenyl] pyrazole (Example C-83) (300 mg, 0.7 mmol) in 50 mL of acetone was added 1 mL of AcOH and NaBH(OAc), (15 g, 70.8 mmol). The mixture was warmed to reflux and was stirred for 5 days. The reaction mixture was poured onto 100 mL of 2.5 N NaOH and was extracted with ethyl acetate (2 x 100 mL). The extracts were combined and washed with brine (1 x 100 mL). The organic phase was dried over Na₂SO₄, filtered, and concentrated to afford the title compound: MS (M+H): 415 (base peak).

Example C-86

5-(4-PIPERIDYL)-4-(4-PYRIDYL)-3-[3-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

By following the method of Example C-1 and substituting methyl-3-(trifluoromethyl)benzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonylisonipecotyl N-hydroxysuccinimide for N-benyloxycarbonylglycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the

title compound as its hydrochloride salt: MS (M+H): 373 (base peak).the pyrazole C-3 substituent (Cviii). Treatment of the

Example C-87

5-(N-METHYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-[3-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

By following the method of Example C-75 and substituting 5-(4-piperidyl)-4-(4-pyridyl)-3-[3-(trifluoromethyl)phenyl) pyrazole hydrochloride (Example C-86) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 387 (base peak).

Example C-88

5-(4-PIPERIDYL)-4-(4-PYRIDYL)-3-(3-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting methyl-3-chlorobenzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonyl-isonipecotyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 339 (base peak).

Example C-89

5-(N-METHYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(3-CHLOROPHENYL)
PYRAZOLE

By following the method of Example C-75 and substituting 5-(4-piperidyl)-4-(4-pyridyl)-3-(3-chlorophenyl) pyrazole hydrochloride (Example C-88) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 353 (base peak).

Example C-90

5-(3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting N-t-butoxycarbonyl-nipecotyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound as its hydrochloride salt: MS (M+H): 323 (base peak).

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Example C-91

5-(N-METHYL-3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-(3-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole hydrochloride (Example C-90) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 337 (base peak).

Example C-92

5-cis-(4-AMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-chlorobenzoate for ethyl-4-

Example C-93

5-cis-(4-N, N-DIMETYLAMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-92) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 381 (base peak).

Example C-94

5-[cis-4-N-(2-PROPYL)AMINOCYCLOHEXYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

a slurry of 5-cis-(4-aminocyclohexyl)-4-(4-To pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-92) (1.0 q, 2.8 mmol, 1.0 eq) in methylene chloride (28 mL) was added acetone (0.5 mL), acetic acid (0.5 mL) and solid sodium triacetoxyborohydride. The slurry was stirred for 5 h and the volatiles were removed. The residue was partitioned between 2.5 M NaOH (25 mL) and ethyl acetate (25 mL) and the aqueous layer was extracted with ethyl acetate (3 x 25 mL). The combined organic layer was washed with brine (50 mL), dried over MgSO, and evaporated. The residue was triturated with ether to yield the title compound as a white powder: 'HNMR (d.-DMSO) δ 8.56 (d, J = 5.84 Hz, 2H), 7.40 (d, J = 8.26 Hz, 2H), 7.30 (d, J = 8.66 Hz, 2H), 7.18 (d, J = 5.64 Hz, 2H), 2.95 (m, 2H), 2.72 (m, 1H), 1.90 (m, 2H), 1.73 (m, 2H), 1.55 (m, 4H), 1.07 (d, J = 5.64 Hz, 6H). MS (M+H): 395 (base peak).

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Example C-95

5-cis-[4-N-(ACETYL)AMINOCYCLOHEXYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-92) for 5-(4piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 395 (base peak).

Example C-96

5-cis-[4-N-(METHOXYACETYL) AMINOCYCLOHEXYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-

(4-chlorophenyl) pyrazole (Example C-92) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) and methoxy acetyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 425 (base peak).

Example C-97

5-cis-[4-n-(METHYLSULFONYL) AMINOCYCLOHEXYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-92) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) and methylsulfonyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 431 (base peak).

Example C-98

5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-Fluorophenyl) pyrazole

By following the method of Example C-1 and substituting N-t-butoxycarbonyl-cis-4-aminocyclohexanoyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 337 (base peak).

Example C-99

5-(cis-4-N, N-DIMETHYLAMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-98) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 365 (base peak).

Example C-100

5-cis-[4-N-(2-PROPYL)AMINOCYCLOHEXYL]-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-94 and substituting cis-5-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-98) for 5-(cis-4-n-(2-propyl)aminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-92) the title compound was prepared: MS (M+H): 379 (base peak).

Example C-101

5-cis-(4-AMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-[4-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-(trifluoromethyl)benzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonyl-cis-4-aminocyclohexanoyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 387 (base peak).

Example C-102

5-cis-(4-N, N-DIMETHYLAMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-[4-(TRIFLUOROMETHYL) PHENYL] PYRAZOLE

By following the method of Example C-75 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-[4-(trifluoromethyl)phenyl] pyrazole (Example C-101) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 415 (base peak).

Example C-103

5-cis-(4-AMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-[3-(TRIFLUOROMETHYL)PHENYL] PYRAZOLE

By following the method of Example C-1 and substituting methyl-3-(trifluoromethyl)benzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonyl-cis-4aminocyclohexanoyl N-hydroxysuccinimide for Nbenyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 387 (base peak).

Example C-104

5-cis-(4-N, N-DIMETHYLAMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-[3-(TRIFLUOROMETHYL) PHENYL] PYRAZOLE

By following the method of Example C-75 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(3-(trifluoromethyl)phenyl) pyrazole (Example C-103) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 415 (base peak).

Example C-105

5-cis-(4-AMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-(3-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting methyl-3-chlorobenzoate for ethyl-4-

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fluorobenzoate and N-t-butoxycarbonyl-cis-4-aminocyclohexanoyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 353 (base peak).

Example C-106

5-cis-(4-N, N-DIMETHYLAMINOCYCLOHEXYL)-4-(4-PYRIDYL)-3-(3-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-cis-(4-aminocyclohexyl)-4-(4-pyridyl)-3-(3-chlorophenyl) pyrazole hydrochloride (Example C-105) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 381 (base peak).

Example C-107

5-(N-ACETIMIDO-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

To a suspension of 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-2) (0.11 g, 0.35 mmol) in 2 mL EtOH was added ethyl acetamidate hydrochloride (0.065 g, 0.53 mmol) and the mixture was refluxed for 30 minutes. The solution was left at 5-10 °C for 16 h and filtered to obtain the title compound as a white solid: MS (M+H): 364 (base peak).

Example C-108

5-(N-CARBOXAMIDINO-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

To a stirred suspension of 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (C-2) (1.5 g, 4.7

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mmol) in 47 mL of DMF was added Hunig's base (0.60 g, 4.7 mmol) and pyrazole carboxamide hydrochloride (0.68 q, 4.7 mmol). The slurry was allowed to stir at room temperature for 4 days. The reaction mixture was poured onto 300 mL of ether. The resulting precipitate was filtered to leave the title compound as the hydrochloride salt: MS (M+H): 365 (base peak).

Example C-109

5-(N-CYCLOPROPANOYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting cyclopropanoyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 407 (base peak).

Example C-110

5~[N-(2-FLUORO)BENZOYL-4-PIPERIDYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 2-fluorobenzoyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 461 (base peak).

Example C-111

5-(N-METHYLSULFONYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-2) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-74)

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and methylsulfonyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 401 (base peak).

Example C-112

5-(N-METHOXYACETYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-2) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-74) and methoxy acetyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 395 (base peak).

Example C-113

5-(N-ACETYL-4-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole Example (C-2) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole Example (C-74) the title compound was prepared: MS (M+H): 365 (base peak).

Example C-114

5-[2-(1,1-DIMETHYL)AMINOETHYL]-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting N-t-butoxycarbonyl-2-amino-2,2-dimethylpropanoyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound as the hydrochloride salt: MS (M+H): 327 (base peak).

Example C-115

5-(METHOXYMETHYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-chlorobenzoate for ethyl-4-fluorobenzoate and 2-methoxyacetyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared: MS (M+H): 300 (base peak).

Example C-116

5-(4-AMINOBENZYL)-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL)

PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-chlorobenzoate for ethyl-4-fluorobenzoate and N-t-butoxycarbonyl-4-aminophenyl

acetyl *N*-hydroxysuccinimide for *N*-benyloxycarbonyl-glycinyl *N*-hydroxysuccinimide the title compound was prepared as the *N*-t-butoxycarbonyl protected compound. The deprotection of the *N*-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound as the hydrochloride salt: MS (M+H): 361 (base peak).

Example C-117

5-[4-(N, N-DIMETHYL) AMINOBENZYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-(4-aminobenzyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-116) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 389 (base peak).

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Example C-118

5-[4-(N-ACETYL)AMINOBENZYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-76 and 5-(4-aminobenzyl)-4-(4-pyridyl)-3-(4substituting chlorophenyl) pyrazole (Example C-116) for 5-(4piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 403 (base peak).

Example C-119

5-(N-METHYLAMINOMETHYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

5-(N-formylaminomethyl)-4-(4-pyridyl)-3-(4fluorophenyl) pyrazole. To a suspension of 5-aminomethylWO 00/31063 PCT/US99/26007

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4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-1) (8.04 g, 30 mmol) in 120 mL dichloromethane was added p-nitrophenylformate (6.01 g, 36 mmol) as a solid. The suspension was stirred for 24 h at room temperature and the solvents removed under reduced pressure. The residue was triturated with ether and filtered to obtain the desired 5-(N-formylaminomethyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole derivative as a white solid: MS (M+H): 297 (base peak).

5-(N-methylaminomethyl)-4-(4-pyridyl)-3-(4-To a suspension of fluorophenyl) pyrazole. 5-(Nformylaminomethyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (8.74 g, 29.5 mmol) in 90 mL anhydrous tetrahydrofuran was added a 1.0 M solution of borane in tetrahydrofuran (90 mL, 90 mmol) and the mixture was stirred at room temperature for 24 h. 1 N aqueous hydrochloric acid (100 mL) was then added to this mixture and the solution was refluxed for 5 hours and cooled to room temperature. The solution was extracted with ether (2 x 250 mL) and the pH of the aqueous layer adjusted to 9 by addition of concentrated ammonium hydroxide. The aqueous layers (pH ~ 9) were then extracted with ethyl acetate (4 x 150 mL). The organic extracts were dried over sodium sulfate, filtered and evaporated to dryness under reduced pressure. The residue was triturated with acetonitrile and filtered to obtain the title compound as a white solid: MS (M+H): 283 (base peak).

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Example C-120

5-[N-(2-AMINO-2,2-DIMETHYLACETYL)AMINOMETHYL]-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

5-(N-t-butoxycarbonylaminomethyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole. To a solution of 5aminomethy1-4-(4-pyridy1)-3-(4-fluoropheny1) pyrazole (Example C-1) (0.27 g, 1 mmol) in anhydrous dimethylformamide (4 mL) was added N-tert-butoxycarbonyl aminoisobutyric acid N-hydroxysuccinimide ester (0.33 g, 1.1 mmol) and the mixture stirred at 40 °C for 24 h. The resulting solution was evaporated to dryness under reduced pressure. The residue was dissolved dichloromethane (30 mL) and washed with a saturated solution of sodium bicarbonate (2 x 20 mL) and brine (20 mL). The organic layers were dried over sodium sulfate, filtered and evaporated under reduced pressure to dryness to afford 5-(N-t-butoxycarbonylaminomethyl)-4-(4pyridyl)-3-(4-fluorophenyl) pyrazole as a white solid.

5-(N-(2-amino-2,2-dimethylacetyl)aminomethyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole. To a solution of the above compound in acetonitrile (2 mL) was added 1 mL of a 4.0 M solution of hydrochloric acid in dioxane. The

reaction mixture was stirred at room temperature for 6 hours. The suspension was evaporated to dryness under reduced pressure. The resulting residue was stirred in acetonitrile (5 mL), filtered and dried in a vacuum dessicator to afford the title compound as a hydrochloride salt: MS (M+H): 354 (base peak).

Example C-121

5-[N-(2-AMINO-2,2-DIMETHYLACETYL) AMINOMETHYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-120 and substituting 5-aminomethyl-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (Example C-15) for 5-aminomethyl-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-1) the title compound was prepared: MS (M+H): 370 (base peak).

Example C-122

5-[4-N-(2-DIMETHYLAMINOACETYL)PIPERIDYL]-4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

To a solution of N,N-dimethylglycine hydrochloride (0.28 g, 2 mmol) in dimethylformamide (4 mL) was added hydroxybenzotriazole (0.27 2 g, mmol), diisopropylethyl amine (0.7 mL, 4 mmol) and polymer supported ethyl carbodimide (Example B-49) (1 g, To this solution after 30 minutes at room temperature was added 5-(4-piperidy1)-4-(4-pyridy1)-3-(4chlorophenyl) pyrazole hydrochloride (Example C-74), 0.41 g, 1 mmol). The suspension was agitated on a labtop orbital shaker for 24 h. The suspension was filtered, washed with dimethylformamide (2 x 5 mL) and the filtrates evaporated under high pressure. The residue was dissolved in dichloromethane (30 mL), washed with a saturated solution of sodium bicarbonate (50 mL) and brine (50 mL). The organic layers were dried over sodium sulfate, filtered and evaporated under high vacuum to afford the title compound as a white solid: MS (M+H): 424 (base peak).

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Example C-123

(S)-5-(2-PYROLIDINYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting (S)-N-t-butoxycarbonyl-prolinyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 309 (base peak).

Example C-124

(S) -5-(N-METHYL-2-PYROLIDINYL) -4-(4-PYRIDYL) -3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting (S)-5-(2-pyrolidinyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-123) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 323 (base peak).

Example C-125

(R) -5-(2-PYROLIDINYL) -4-(4-PYRIDYL) -3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting (R)-N-t-butoxycarbonyl-prolinyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 309 (base peak).

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Example C-126

(R)-5-(N-METHYL-2-PYROLIDINYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting (R)-5-(2-pyrolidinyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-125) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 323 (base peak).

Example C-127

(R)-5-(3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting (R)-N-t-butoxycarbonyl-nipecotyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-

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hydroxysuccinimide the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound: MS (M+H): 323 (base peak).

Example C-128

(R)-5-(N-METHYL-3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting (R)-5-(3-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (Example C-125) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole hydrochloride (Example C-74) the title compound was prepared: MS (M+H): 337 (base peak).

Example C-129

2,2-DIMETHYL-4-[4-(4-PYRIDYL)-3-(4-CHLOROPHENYL) PYRAZOLYL] BUTYRIC ACID

By following the method of Example C-1 and substituting methyl-4-chlorobenzoate for ethyl-4-fluorobenzoate and 2,2-dimethyl glutaric anhydride for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide the title compound was prepared: MS (M+H): 370 (base peak).

Example C-130

4-[4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLYL] BUTYRIC ACID

By following the method of Example C-1 and substituting glutaric anhydride for N-benzyloxycarbonylglycinyl N-hydroxysuccinimide the title compound was prepared: MS (M+H): 326 (base peak).

Example C-131

4-[4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLYL] BUTYRAMIDE

Methvl 4-(4-(4-pyridyl)-3-(4-fluorophenyl)pyrazolyl) butyrate. To a solution of 4-(4-(4-pyridyl)-3-(4-fluorophenyl) pyrazolyl) butyric acid (Example C-130) (40 g, 123 mmol) in 650 mL of MeOH was added 20 mL of concentrated H,SO,. The solution was stirred overnight at room temperature. The solution was concentrated and diluted with 200 mL of water. The solution was cooled with an ice/water bath and to the solution was added 150 mL of saturated NaHCO,. The solution was neutralized further with 50% NaOH to pH 7. The resulting slurry was extracted with CH,Cl, (3 x 250 mL). The combined extracts were washed with water (1 x 300 mL) and saturated NaHCO, (1 x 500 mL). The organic phase was dried over Na,SO, filtered and concentrated to afford methyl 4-(4-(4pyridyl)-3-(4-fluorophenyl) pyrazolyl) butyrate: (M+H): 340 (base peak).

4-(4-(4-pyridyl)-3-(4-fluorophenyl) pyrazolyl)
butyramide. A solution of methyl 4-(4-(4-pyridyl)-3-(4fluorophenyl) pyrazolyl) butyrate (39 g, 120 mmol) in 600
mL of MeOH was saturated with NH,. The solution was

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periodically treated with additional NH, over a 24 h period. The solution was degassed with a stream of nitrogen and the solution was concentrated to leave a yellow solid. The solid was slurried in ether and filtered to leave the title compound: MS (M+H): 325 (base peak).

Example C-132

5-[4-(1-HYDROXY)BUTYL]-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

A stirred suspension of 4-(4-(4-pyridyl)-3-(4-fluorophenyl) pyrazolyl) butyric acid (Example C-130) (2 g, 6.15 mmol) in 100 ml of anhydrous ether was cooled to 0 °C under nitrogen. Lithium aluminum hydride (467 mg, 12.3 mmol) was added to this suspension slowly. After the addition was complete, the mixture was warmed to room temperature and stirred for additional 2 h. The reaction was quenched slowly with 1N KHSO, (80 ml). The mixture was transferred to a separatory funnel and the aqueous layer was removed. The aqueous layer was then made basic with K,CO, (pH 8). The aqueous solution was extracted with ethyl acetate (2 x 100 mL). The combined ethyl acetate extracts were washed with water (1 x 100

mL), dried over MgSO, filtered and concentrated to give the title compound: MS (M+H): 312 (base peak).

Example C-133

5-[4-(1,1-DIMETHYL-1-HYDROXY)BUTYL]-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

A solution of 4-(4-(4-pyridyl)-3-(4-fluorophenyl) pyrazolyl) butyric acid (Example C-130) (200 mg, 0.615 mmol) in 50 ml of MeOH was treated with 10 ml of 4 N HCl/dioxane. The reaction mixture was stirred for 5 hours and evaporated to dryness. To this residue was added 15 ml of 1N methyl magnesium bromide in butyl ether and 5 ml of anhydrous THF. The reaction was heated to reflux under nitrogen for 64 h.

The reaction was quenched with 20 ml of saturated ammonium chloride. This mixture was transferred to a separatory funnel and was extracted with 100 ml ethyl acetate (2 x 100 mL). The combined ethyl acetate extracts were washed with water (1 x 100 mL), dried over MgSO₄, filtered and concentrated to afford a crude oil. The crude oil was subjected to column chromatography by using 3.5 % MeOH/CH₂Cl₂ followed by 6 % MeOH/CH₂Cl₂ to give the title compound: MS (M+H): 340 (base peak).

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Example C-134

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5-(4-(1-AMINO)BUTYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

suspension of To 4-(4-(4-pyridyl)-3-(4fluorophenyl) pyrazolyl) butyramide (Example C-131) (2 g, 6.2 mmol) in 100 ml of anhydrous ether was added lithium aluminum hydride (467 mg, 12.3 mmol). After the addition was complete, the mixture was warmed to room temperature and stirred for additional 2 h. The reaction was quenched with 20 mL of ethyl acetate and was poured onto 100 mL of 2.5 N NaOH. The mixture was extracted with ethyl acetate (3 x 50 mL). The combined extracts were washed with brine (1 x 100 mL), dried over Na,SO,, filtered and concentrated to afford the title compound: MS (M+H): 311 (base peak).

Example C-135

4-(4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLYL) PROPIONIC ACID

N-NH OH

By following the method of Example C-1 and substituting succinic anhydride for N-benyloxycarbonylglycinyl N-hydroxysuccinimide the title compound was prepared: MS (M+H): 312 (base peak).

Example C-136

5-(4-PIPERIDYL)-4-(4-PYRIMIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-1 and substituting methyl-4-chlorobenzoate for ethyl-4-fluorobenzoate, N-t-butoxycarbonyl-isonipecotyl N-hydroxysuccinimide for N-benyloxycarbonyl-glycinyl N-hydroxysuccinimide and 4-methylpyrimidine for 4-picoline

the title compound was prepared as the N-t-butoxycarbonyl protected compound. The deprotection of the N-t-butoxycarbonyl intermediate was accomplished with 4 N HCl in dioxane to afford the title compound as the hydrochloride salt: ¹H NMR (CDCl₃) δ 9.2 (s, 1 H), 8.48 (d, J = 5.19 Hz, 1 H), 7.31 (m, 4 H), 6.94 (d, J = 4.79 Hz, 1 H), (3.69 (m, 3 H), 3.12 (m, 2 H), 2.3 (m, 3 H), 1.24 (m, 2 H). MS (M+H): 340 (base peak).

Example C-137

5-(N-METHYL-4-PIPERIDYL)-4-(4-PYRIMIDYL)-3-(4-CHLOROPHENYL) PYRAZOLE

By following the method of Example C-75 and substituting 5-(4-piperidy1)-4-(4-pyrimidy1)-3-(4-chloropheny1) pyrazole (Example C-136) for 5-(4-piperidy1)-4-(4-pyridy1)-3-(4-chloropheny1) pyrazole hydrochloride (Example C-74) the title compound was prepared: 'H NMR (CDCl₁) δ 9.2 (d, J = 1.2 Hz, 1 H), 8.48 (d, J = 5.59 Hz, 1 H), 7.31 (m, 4 H), 6.95 (dd, J= 1.2, 5.6 Hz, 1 H), 3.39 (m, 1 H), 3.03 (d, J = 11.6 Hz, 2 H), 2.38 (s, 3 H), 2.06 (m, 4 H), 1.24 (m, 2 H). MS (M+H): 354 (base peak).

Example C-138

5-(N-ACETYL-3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-(3-piperidy1)-4-(4-pyridy1)-3-(4-fluoropheny1) pyrazole (C-90) for 5-(4-piperidy1)-4-(4-pyridy1)-3-(4-chloropheny1) pyrazole (C-74) the title compound was prepared: MS (M+H): 365 (base peak).

Example C-139

5-(N-METHOXYACETYL-3-PIPERIDYL)-4-(4-PYRIDYL)-3-(4-FLUOROPHENYL) PYRAZOLE

By following the method of Example C-76 and substituting 5-(3-piperidyl)-4-(4-pyridyl)-3-(4-fluorophenyl) pyrazole (C-90) for 5-(4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl) pyrazole (C-74) and methoxy

acetyl chloride for acetyl chloride the title compound was prepared: MS (M+H): 395 (base peak).

Additional compounds of the present invention which could be prepared using one or more of the reaction schemes set forth in this application include, but are not limited to, the following:

Example C-140

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-thiomethyl)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-141

5-(4-piperidinyl)-4-[4-(2-thiomethyl)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

5-(4-N-methylpiperidinyl)-4-[4-(2-thiomethyl)pyrimidinyl]-3-4-(chlorophenyl)pyrazole

Example C-143

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-methanesulfonyl)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-144

5-(4-piperidinyl)-4-[4-(2-methanesulfonyl)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

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Example C-145

5-(4-N-methylpiperidinyl)-4-[4-(2-methanesulfonyl)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-146

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-amino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-147

5-(4-piperidinyl)-4-[4-(2-amino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

5-(4-N-methylpiperidinyl)-4-[4-(2-amino)pyrimidinyl]-3(4-chlorophenyl)pyrazole

Example C-149

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-methylamino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-150

5-(4-piperidinyl)-4-[4-(2-methylamino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

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Example C-151

5-(4-N-methylpiperidinyl)-4-[4-(2-

methylamino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-152

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-isopropylamino)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-153

5-(4-piperidiny1)-4-[4-(2-isopropylamino)pyrimidiny1]-3(4-chloropheny1)pyrazole

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Example C-154

5-(4-N-methylpiperidinyl)-4-[4-(2-

isopropylamino)pyrimidiny1]-3-(4-chlorophenyl)pyrazole

Example C-155

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-(2-methoxyethylamino))pyrimidinyl]-3-(4-

chlorophenyl)pyrazole

Example C-156

5-(4-piperidiny1)-4-[4-(2-(2-

methoxyethylamino))pyrimidinyl]-3-(4-

chlorophenyl)pyrazole

5-(4-N-methylpiperidinyl)-4-[4-(2-(2-methoxyethylamino))pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-158

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-methoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-159

5-(4-piperidiny1)-4-[4-(2-methoxy)pyrimidiny1]-3-(4-chloropheny1)pyrazole

Example C-160

5-(4-N-methylpiperidinyl)-4-[4-(2-methoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-161

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-isopropoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-162

5-(4-piperidinyl)-4-[4-(2-isopropoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-163

5-(4-N-methylpiperidinyl)-4-[4-(2-isopropoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-164

5-(4-N-t-butoxycarbonylpiperidinyl)-4-[4-(2-(2-N,N-dimethylamino)ethoxy)pyrimidinyl]-3-(4-

chlorophenyl)pyrazole

Example C-165

5-(4-piperidinyl)-4-[4-(2-(2-N,N-dimethylamino)ethoxy)pyrimidinyl]-3-(4-

chlorophenyl)pyrazole

Example C-166

5-(4-N-methylpiperidinyl)-4-[4-(2-(2-N,N-dimethylamino)ethoxy)pyrimidinyl]-3-(4-chlorophenyl)pyrazole

Example C-167

5-(N-acetylhydroxylimido-4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-168

5-(N-benzylhydroxylimido-4-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

5-(N-phenylacethydroxylimido-4-piperidyl)-4-(4-pyridyl)3-(4-chlorophenyl)pyrazole

Example C-170

5-[N-methyl-4-(3,4-dehydro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-171

5-[N-isopropyl-4-(3,4-dehydro)piperidyl]-4-(4-pyridyl)-3(4-chlorophenyl)pyrazole

5-[N-benzyl-4-(3,4-dehydro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-173

5-[N-methyl-4-(4-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-174

5-[N-methyl-4-(4-hydroxy)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-175

5-[N-methyl-4-(4-methoxy)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-176

5-[N-methyl-4-(2,5-tetramethyl-4-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-177

5-[N-methyl-4-(2,5-tetramethyl-4-hydroxy)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-178

5-[N-methyl-4-(2,5-tetramethyl-4-methoxy)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-179

5-[4-(3-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-180

5-[4-(N-methyl-3-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-181

5-[4-(N-isopropyl-3-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-182

5-[4-(N-benzyl-3-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-183

5-[4-(N-acetyl-3-fluoro)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-184

5-[4-(2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-185

5-[4-(N-methyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-186

5-[4-(N-isopropyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-187

5-[4-(N-benzyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-188

5-[4-(N-acetyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-189

5-[5-(2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-190

5-[5-(N-methy1-2-oxo)piperidy1]-4-(4-pyridy1)-3-(4-chloropheny1)pyrazole

Example C-191

5-[5-(N-isopropyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-192

5-[5-(N-benzyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-193

5-[5-(N-acetyl-2-oxo)piperidyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-194

5-(N-acethydroxylimido-3-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

5-(N-benzhydroxylimido-3-piperidyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-196

5-(N-phenacethydroxylimido-3-piperidyl)-4-(4-pyridyl)-3(4-chlorophenyl)pyrazole

Example C-197

5-(2-morpholinyl)-4-(4-pyridyl)-3-(4chlorophenyl)pyrazole

5-(N-methyl-2-morpholinyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-199

5-(N-isopropyl-2-morpholinyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-200

5-(N-benzyl-2-morpholinyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

5-(N-acetyl-2-morpholinyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-202

5-[trans-4-(N-t-butoxycarbonylamino)methylcyclohexyl]-4(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-203

5-(trans-4-aminomethylcyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-204

5-[trans-4-(N-isopropylamino)methylcyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-205

5-[trans-4-(N, N-dimethylamino)methylcyclohexyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-206

5-[trans-4-(N-acetylamino)methylcyclohexyl)]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-207

5-[trans-4-(N-t-butoxycarbonylamino)cyclohexyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-208

5-(trans-4-aminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-209

5-[trans-4-(N,N-dimethylamino)cyclohexyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-210

5-[trans-4-(N-isopropylamino)cyclohexyl)-4-(4-pyridyl)-3(4-chlorophenyl)pyrazole

Example C-211

5-[trans-4-(N-acetylamino)cyclohexyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-212

5-[cis-4-(N-t-butoxycarbonyl)methylaminocyclohexyl)]-4(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-213

5-(cis-4-methylaminocyclohexyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-214

5-[cis-4-(N,N-dimethyl)methylaminocyclohexyl)]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-215

5-[cis-4-(N-isopropyl)methylaminocyclohexyl)]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-216

5-[cis-4-(N-acetyl)methylaminocyclohexyl)]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-217

5-[3-(1,1-dimethyl-1-(N-t-butoxycarbonylamino)propyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-218

5-[3-(1,1-dimethyl-1-amino)propyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

1000

Example C-219

5-[3-(1,1-dimethyl-1-(N,N-dimethylamino)propyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-220

5-[3-(1,1-dimethyl-1-(N-isopropylamino)propyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-221

5-[3-(1,1-dimethyl-1-(N-acetylamino)propyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-222

5-[4-(1-carboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-223

5-[4-(1-N-methylcarboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-224

5-[4-(1-N-benzylcarboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-225

5-[3-(1-carboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-226

5-[3-(1-N-methylcarboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-227

5-[3-(1-N-benzylcarboxamidino)benzyl-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-228

5-[3-(N-t-butoxycarbonyl)aminobenzyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-229

5-(3-aminobenzyl)-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-230

5-[3-(N, N-dimethylamino)benzyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

1004

Example C-231

5-[3-(N-isopropylamino)benzyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-232

5-[3-(N-benzylamino)benzyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

1005

Example C-233

5-[3-(N-acetylamino)benzyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-234

5-[4-(2-amino)methylimidazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-235

5-[4-(2-N, N-dimethylamino)methylimidazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

1006

Example C-236

5-[4-(2-N-isopropylamino)methylimidazolyl]-4-(4-pyridyl)3-(4-chlorophenyl)pyrazole

Example C-237

5-[4-(2-N-benzylamino)methylimidazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-238

5-[4-(2-N-acetylamino)methylimidazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-239

5-[4-(2-amino)methyloxazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-240

5-[4-(2-N, N-dimethylamino)methyloxazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-241

5-[4-(2-N-isopropylamino)methyloxazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

100B

Example C-242

5-[4-(2-N-benzylamino)methyloxazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-243

5-[4-(2-N-acetylamino)methyloxazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

Example C-244

5-[4-(2-amino)methylthiazolyl]-4-(4-pyridyl)-3-(4chlorophenyl)pyrazole

Example C-245

5-[4-(2-N, N-dimethylamino)methylthiazolyl]-4-(4-pyridyl)3-(4-chlorophenyl)pyrazole

Example C-246

5-[4-(2-N-isopropylamino)methylthiazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

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Example C-247

5-[4-(2-N-benzylamino)methylthiazolyl]-4-(4-pyridyl)-3(4-chlorophenyl)pyrazole

Example C-248

5-[4-(2-N-acetylamino)methylthiazolyl]-4-(4-pyridyl)-3-(4-chlorophenyl)pyrazole

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Biological data from compounds of Examples B-0001 through B-1573 and of Examples B-2270 through B-2462 are shown in the following tables.

In vitro P38-alpha kinase inhibitory data are shown in the column identified as:

"P38 alpha kinase IC50, uM or % inhib @ conc. (uM)"

10

15

25

30

In vitro whole cell assay for measuring the ability of the compounds to inhibit TNF production in human U937 cells stimulated with LPS are shown in the column identified as:

"U937 Cell IC50, uM or % inhib @ conc., (uM)"

In vivo assessment of the ability of the compounds to inhibit LPS-stimulated TNF release in the mouse is shown in the column identified as:

"Mouse LPS Model, % TNF inhib @ dose @ predose time"
wherein in the dose is milligram per kilogram (mpk)
administered by oral gavage and the predose time
indicates the number of hours before LPS challenge when
the compound is administered.

In vivo assessment of the ability of the compounds to inhibit LPS-stimulated TNF release in the rat is shown in the column identified as:

"Rat LPS Model, % TNF inhib @ dose @ predose time"

wherein in the dose is milligram per kilogram (mpk)

administered by oral gavage and the predose time

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indicates the number of hours before LPS challenge when the compound is administered.

	P38 alpha kinase IC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or % inhlb@conc. (uM)	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
Example#				
B-0001	53.0%@1.0uM	40.0% @1.0uM		
B-0002	71.0%@1.0uM	28.0%@10.0uM		
B-0003	70.0%@1.0uM	76.0% 10.0uM		
B-0004	80.0%@1.0uM	4.61uM		
B-0005	95.0%@1.0uM	2.97uM		
B-0006	82.0%@1.0uM	80%@10.0uM		
B-0007	74.0%@1.0uM	85.0%@10.0uM		
B-0008	42.0%@1.0uM	65.0%@10.0uM		
B-0009	0.04 uM	0.72uM		
B-0010	0.52 uM	0.65uM		
B-0011	0.03 uM	4.47uM		
B-0012	30.0%@1.0uM	44.0% @1.0uM		
B-0013	70.0%@1.0uM	84.0%@10.0uM		
B-0014	79.0%@1.0uM	80.0%@10.0uM		
B-0015	82.0%@1.0uM	80.0%@10.0uM		
B-0016	94.0%@1.0uM	3,98uM		
B-0017	56.0%@1.0uM	79.0%@10.0uM		
B-0018	60.0%@1.0uM	59.0%@10.0uM		
B-0019	84.0%@1.0uM	100.0%@10.0uM		· · · · · · · · · · · · · · · · · · ·
8-0020	73.0%@1.0uM	81.0%@10.0uM	1	
B-0021	68.0%@1.0uM	76.0%@10.0uM		
B-0022	69.0%@1.0uM	44.0@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0023	90.0%@1.0uM	77.0%@10.0uM		
B-0024	94.0%@1.0uM	52.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0025	89.0%@1.0uM	79.0%@10.0uM		
8-0026	96.0%@1.0uM	3.27uM		
B-0027	94.0%@1.0uM	11.0uM		· · · · · · · · · · · · · · · · · · ·
B-0028	69.0%@1.0uM	45.0%@10.0uM		
B-0029	91.0%@1.0uM	58.0%@10.0uM		
B-0030	92.0%@1.0uM	75.0%@10.0uM		·- · · · · · · · · · · · · · · · · · ·
B-0031	94.0%@1.0uM	100.0%@10.0uM		
B-0032	94.0%@1.0uM	78.0%@10.0uM		
B-0033	97.0%@1.0uM	10.0uM		
B-0034	95.0%@1.0uM	10.0uM	· · · · · · · · · · · · · · · · · · ·	
B-0035	94.0%@1.0uM	10.0uM		
B-0036	92.0%@1.0uM	8.24uM		
B-0037	91.0%@1.0uM	86.0%@10.0uM	,	
B-0038	71.0%@1.0uM	84.0%@10.0uM		
B-0039	89.0%@1.0uM	72.0%@10.0uM		-
B-0040	93.0%@1.0uM	2.3uM		
B-0041	65.0%@1.0uM	66.0%@10.0uM		
B-0042	94.0%@1.0uM	2.76uM		***************************************

IC50,uM or % Inhib@conc. (uM)			I		
Inhib@conc. (uM)					Rat LPS Model %
Example# B-0043 0.22 uM 0.54uM B-0044 0.14 uM 0.19uM B-0045 94.0%@1.0uM 1.01uM B-0046 96.0%@1.0uM 54.0%@1.0uM B-0047 94.0%@1.0uM 74.0%@10.0uM B-0048 94.0%@1.0uM 76.0%@10.0uM B-0049 88%@1.0uM 33.0%@1.0uM B-0050 73%@1.0uM 34.0%@1.0uM B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1.0uM 34.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM 31.0uM 31.0uM B-0055 93%@1.0uM 21.0%@1.0uM B-0055 96%@1.0uM 21.0%@1.0uM B-0056 96%@1.0uM 22.0%@1.0uM B-0058 79%@1.0uM 28.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 4.0%@1.0uM B-0063 96%@1.0uM 4.0%@1.0uM B-0066 94%@1.0uM 4.0%@1.0uM B-0066 94%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 27.0%@1.0uM B-0067 91%@1.0uM 27.0%@1.0uM 27.0%@1.0uM B-0067 91%@1.0uM 27.0%@1.0uM 27.0%@1.0uM B-0067 91%@1.0uM 27.0%@1.0uM 27.0%@1.0uM B-0067 91%@1.0uM 27.0%@1.0uM 27.0%@1.0uM 27.0%@1.0uM 37.0%@1.0uM 37.00M@1.0uM 37.00M@1.0uM 37.00M@1.0uM 37.00M					
B-0044	Example#		` '		
B-0045 94.0%@1.0uM 1.01uM B-0046 96.0%@1.0uM 54.0%@1.0uM B-0047 94.0%@1.0uM 74.0%@10.0uM B-0048 94.0%@1.0uM 76.0%@10.0uM B-0049 88%@1.0uM 33.0%@1.0uM B-0050 73%@1.0uM 34.0%@1.0uM B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1.0uM 15.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM 21.0uM B-0056 96%@1.0uM 21.00@1.0uM B-0057 96%@1.0uM 22.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 22.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 27.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 40.0%@1.0uM B-0066 94%@1.0uM 21.0%@1.0uM B-0067 91%@1.0uM 22.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0060 92%@1.0uM 37.0%@1.0uM B-0060 92%@1.0uM 37.0%@1.0uM B-0060 92%@1.0uM 37.0%@1.0uM B-0060 92%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 37.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0043	0.22 uM	0.54uM		
B-0046 96.0%@1.0uM 54.0%@1.0uM B-0047 94.0%@1.0uM 74.0%@10.0uM B-0048 94.0%@1.0uM 76.0%@10.0uM B-0049 88%@1.0uM 33.0%@1.0uM B-0050 73%@1.0uM 34.0%@1.0uM B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1.0uM 34.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM >1.0uM B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 22.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 28.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 28.0%@1.0uM B-0068 72%@1.0uM 28.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 37.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0044	0.14 uM	0.19uM		
B-0047 94.0%@1.0uM 74.0%@10.0uM B-0048 94.0%@1.0uM 76.0%@10.0uM B-0049 88%@1.0uM 33.0%@1.0uM B-0050 73%@1.0uM 34.0%@1.0uM B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1.0uM 15.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM >1.0uM B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 22.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 28.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 28.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0045	94.0%@1.0uM	1.01uM		
B-0048 94,0%@1,0uM 76,0%@10,0uM B-0049 88%@1,0uM 33,0%@1,0uM B-0050 73%@1,0uM 34,0%@1,0uM B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1,0uM 15,0%@1,0uM B-0053 95%@1,0uM 34,0%@1,0uM B-0054 90%@1,0uM 30,0%@1,0uM B-0055 93%@1,0uM >1,0uM B-0056 96%@1,0uM 21,0%@1,0uM B-0057 96%@1,0uM 29,0%@1,0uM B-0058 79%@1,0uM 18,0%@1,0uM B-0059 83%@1,0uM 35,0%@1,0uM B-0060 73%@1,0uM 22,0%@1,0uM B-0061 62%@1,0uM 27,0%@1,0uM B-0062 94%@1,0uM 36,0%@1,0uM B-0063 96%@1,0uM 40,0%@1,0uM B-0064 90%@1,0uM 40,0%@1,0uM B-0065 83%@1,0uM 21,0%@1,0uM B-0066 94%@1,0uM 21,0%@1,0uM B-0067 91%@1,0uM 21,0%@1,0uM B-0068 72%@1,0uM 22,0%@1,0uM B-0069 96%@1,0uM 21,0%@1,0uM B-0069 96%@1,0uM 37,0%@1,0uM B-0069 96%@1,0uM 37,0%@1,0uM B-0069 96%@1,0uM 37,0%@1,0uM B-0069 96%@1,0uM 37,0%@1,0uM B-0070 92%@1,0uM 31,0%@1,0uM B-0071 86%@1,0uM 31,0%@1,0uM B-0072 77%@1,0uM 32,0%@1,0uM B-0073 91%@1,0uM 32,0%@1,0uM B-0073 91%@1,0uM 32,0%@1,0uM	B-0046	96.0%@1.0uM	54.0%@1.0uM		
B-049	B-0047	94.0%@1.0uM	74.0%@10.0uM		
B-0050	B-0048	94.0%@1.0uM	76.0%@10.0uM		
B-0051 3.3uM 2.15uM 47%@100mpk@-6h 79%@3mpk@-4h B-0052 92%@1.0uM 15.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 31.0uM 31.0uM S-0055 93%@1.0uM 21.0%@1.0uM S-0057 96%@1.0uM 29.0%@1.0uM S-0058 79%@1.0uM 18.0%@1.0uM S-0059 83%@1.0uM 35.0%@1.0uM S-0059 83%@1.0uM 22.0%@1.0uM S-0061 62%@1.0uM 27.0%@1.0uM S-0062 94%@1.0uM 36.0%@1.0uM S-0063 96%@1.0uM 40.0%@1.0uM S-0064 90%@1.0uM 40.0%@1.0uM S-0065 83%@1.0uM 21.0%@1.0uM S-0066 94%@1.0uM 21.0%@1.0uM S-0066 94%@1.0uM 21.0%@1.0uM S-0066 94%@1.0uM 21.0%@1.0uM S-0067 91%@1.0uM 22.0%@1.0uM S-0068 72%@1.0uM 22.0%@1.0uM S-0069 96%@1.0uM 37.0%@1.0uM S-0069 96%@1.0uM 37.0%@1.0uM S-0069 96%@1.0uM 30.0%@1.0uM S-0070 92%@1.0uM 31.0%@1.0uM S-0071 86%@1.0uM 31.0%@1.0uM S-0072 77%@1.0uM 32.0%@1.0uM S-0073 91%@1.0uM S-0078	B-0049	88%@1.0uM	33.0%@1.0uM		
B-0052 92%@1.0uM 15.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM >1.0uM B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 29.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 21.0%@1.0uM B-0067 91%@1.0uM 22.0%@1.0uM B-0068 72%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0050	73%@1.0uM	34.0%@1.0uM		
B-0052 92%@1.0uM 15.0%@1.0uM B-0053 95%@1.0uM 34.0%@1.0uM 30.0%@1.0uM B-0054 90%@1.0uM 31.0uM 31.0	B-0051	3.3uM	2.15uM	47%@100mpk@-6h	79%@3mpk@-4h
B-0054 90%@1.0uM 30.0%@1.0uM B-0055 93%@1.0uM >1.0uM B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 29.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 22.0%@1.0uM B-0067 91%@1.0uM 22.0%@1.0uM B-0068 72%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0052	92%@1.0uM	15.0%@1.0uM		
B-0055 93%@1.0uM >1.0uM B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 29.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 40.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 21.0%@1.0uM B-0067 91%@1.0uM 22.0%@1.0uM B-0068 72%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0053	95%@1.0uM	34.0%@1.0uM		
B-0056 96%@1.0uM 21.0%@1.0uM B-0057 96%@1.0uM 29.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 22.0%@1.0uM B-0067 91%@1.0uM 28.0%@1.0uM B-0068 72%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0054	90%@1.0uM	30.0%@1.0uM		
B-0057 96%@1.0uM 29.0%@1.0uM B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 37.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 32.0%@1.0uM	B-0055	93%@1.0uM	>1.0uM		
B-0058 79%@1.0uM 18.0%@1.0uM B-0059 83%@1.0uM 35.0%@1.0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0065 94%@1.0uM 22.0%@1.0uM B-0066 94%@1.0uM 22.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0056	96%@1.0uM	21.0%@1.0uM		
B-0059 83%@1,0uM 35.0%@1,0uM B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1,0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0065 94%@1.0uM 22.0%@1.0uM B-0066 94%@1.0uM 22.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0057	96%@1.0uM	29.0%@1.0uM		************
B-0060 73%@1.0uM 22.0%@1.0uM B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 22.0%@1.0uM	B-0058	79%@1.0uM	18.0%@1.0uM		
B-0061 62%@1.0uM 27.0%@1.0uM B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0059	83%@1.0uM	35.0%@1.0uM		
B-0062 94%@1.0uM 36.0%@1.0uM B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0060	73%@1.0uM	22.0%@1.0uM		
B-0063 96%@1.0uM 40.0%@1.0uM B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0061	62%@1.0uM	27.0%@1.0uM		
B-0064 90%@1.0uM 4.0%@1.0uM B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0062	94%@1.0uM	36.0%@1.0uM		
B-0065 83%@1.0uM 21.0%@1.0uM B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0063	96%@1.0uM	40.0%@1.0uM		
B-0066 94%@1.0uM 28.0%@1.0uM B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0064	90%@1.0uM	4.0%@1.0uM		
B-0067 91%@1.0uM 1.0%@1.0uM B-0068 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0065	83%@1.0uM	21.0%@1.0uM		
B-0058 72%@1.0uM 22.0%@1.0uM B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0066	94%@1.0uM	28.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0069 96%@1.0uM 37.0%@1.0uM B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0067	91%@1.0uM	1.0%@1.0uM		····
B-0070 92%@1.0uM 30.0%@1.0uM B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0068	72%@1.0uM	22.0%@1.0uM		
B-0071 86%@1.0uM 31.0%@1.0uM B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0069	96%@1.0uM	37.0%@1.0uM		
B-0072 77%@1.0uM 32.0%@1.0uM B-0073 91%@1.0uM 24.0%@1.0uM	B-0070	92%@1.0uM	30.0%@1.0uM		
B-0073 91%@1.0uM 24.0%@1.0uM	B-0071	86%@1.0uM	31.0%@1.0uM		
	B-0072	77%@1.0uM	32.0%@1.0uM		
B-0074 92%@1 0ulk 42.0%@1 0ulk	B-0073	91%@1.0uM	24.0%@1.0uM		
1.0017 32.08 1.00M 42.0%8 1.00M	B-0074	92%@1.0uM	42.0%@1.0uM		
B-0075 91%@1.0uM 35.0%@1.0uM	B-0075	91%@1.0uM	35.0%@1.0uM		*
B-0076 58%@1.0uM 21.0%@1.0uM	B-0076	58%@1.0uM	21.0%@1.0uM		
B-0077 0.8uM 10.0uM	B-0077	0.8uM	10.0uM		
B-0078 80%@1.0uM 20.0%@1.0uM	B-0078	80%@1.0uM	20.0%@1.0uM		
B-0079 93%@1.0uM 13.0%@1.0uM	B-0079	93%@1.0uM	13.0%@1.0uM		
B-0080 73%@1.0uM 73.0%@1.0uM	B-0080	73%@1.0uM	73.0%@1.0uM		
B-0081 92%@1.0uM 13.0%@1.0uM	B-0081	92%@1.0uM	13.0%@1.0uM		
B-0082 47%@1.0uM 27.0%@1.0uM	B-0082	47%@1.0uM	27.0%@1.0uM		
B-0083 0.22uM 6.51uM	B-0083	0.22uM	****		
B-0084 56%@1.0uM 30.0%@1.0uM	B-0084	56%@1.0uM	30.0%@1.0uM		

Example#	P38 alpha kinase IC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or % inhib@conc. (uM)	TNF inhib @ dose	
B-0085	83%@1.0uM	21.0%@1.0uM		
B-0086	91%@1.0uM	37.0%@1.0uM		
B-0087	0.55uM	2.26uM	38%@30mpk@-6h	
B-0088	96%@1.0uM	9.0%@1.0uM	CONCOUNTRE ON	
B-0089	0.04uM	3.33uM		
B-0090	98%@1.0uM	52.0%@1.0uM	 	
B-0091	96%@1.0uM	40.0%@1.0uM		
B-0092	97%@1.0uM	34.0%@1.0uM		
B-0093	3.18 uM	1.25uM	30%@30mpk@-6h	
B-0094	96%@1.0uM	52.0%@1.0uM	Control Compare on	· · · · · · · · · · · · · · · · · · ·
B-0095	98%@1.0uM	38.0%@1.0uM		
B-0096	91%@1.0uM	22.0%@1.0uM		
B-0097	72.0%@10.0uM	38.0%@1.0uM		
B-0098	66.0%@10.0uM	12.0%@1.0uM		
B-0099	43.0% @1.0uM	>1.0uM		
3-0100	75.0% @1.0uM	5,0uM		
B-0101	71.0% @ 1.0uM	2.11uM		
3-0102	81.0%@1.0uM	15.0%@1.0uM		
B-0103	71.0%@1.0uM	6.0%@1.0uM		
3-0104	56.0% @1.0uM	2.78uM		
3-0105	78.0%@1.0uM	5.0uM		
3-0106	62.0%@1.0uM	5.0uM		
3-0107	0.27uM	5.0uM		
3-0108	61.0%@1.0uM	4.85uM		
3-0109	45.0%@1.0uM	19.0%@1.0uM		
3-0110	66.0%@1.0uM	13.0%@1.0uM		
3-0111	57.0%@1.0uM	>1.0uM		
3-0112	97.0%@1.0uM	1.12uM		
3-0113	75.0%@1.0uM	43.0%@1.0uM		
-0114	45.0%@1.0uM	3.92uM		
-0115	47.0%@1.0uM	2.0%@1.0uM		
-0116	73.0%@1.0uM	35.0%@1.0uM		
-0117	0.46 uM	1.78 uM	30%@30mpk@-6h	
-0118	1.18 uM	1.29 uM		
-0119	89.0%@10.0uM	2.78uM		
-0120	Mu 800.0	0.21 uM	77%@100mpk@-6h	70%@3mpk@-4h
-0121	79.0%@1.0uM	1.22uM		· · · · · · · · · · · · · · · · · · ·
-0122	79.0%@10.0uM	2.0%@1.0uM		
-0123	59.0%@1.0uM	>1.0uM		
-0124	73.0%@1.0uM	15.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
-0125	70.0%@10.0uM	17.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
-0126	66.0%@1.0uM	1.57uM		<u> </u>

F				
	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or % inhib@conc. (uM)	TNF inhib @ dose	inhib @dose @predose time
Example#	inhib@conc. (uM)	innib@conc. (um)	@predose time	wpredose time
B-0127	82.0%@1.0uM	0.96uM		
B-0128	78.0%@1.0uM	1.81uM		
B-0129	51.0%@1.0uM	31.0%@1.0uM		
B-0130	69.0%@1.0uM	58.0%@1.0uM		
B-0131	43.0%@1.0uM	46.0%@1.0uM		
B-0132	76.0%@1.0uM	8.0%@1.0uM		
B-0133	51.0%@1.0uM	42.0%@1.0uM		
B-0134	60.0%@1.0uM	2,17uM		
B-0135	78.0%@1.0uM	58.0%@1.0uM		
B-0136	77.0%@1.0uM	44.0%@1.0uM		
B-0137	41.0%@1.0uM	37.0%@1.0uM		
		32.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0138	50.0%@1.0uM			
B-0139	54.0%@10.0uM	17.0%@1.0uM		
B-0140	67%@10.0uM	9.0%@1.0uM		
B-0141	78.0%@1.0uM	10.0%@1.0uM		
B-0142	86.0%@1.0uM	12.0%@1.0uM		
B-0143	42.0% @1.0uM	3.63uM		
B-0144	86.0% @1.0uM	43.0%@1.0uM		
B-0145	54.0% @10.0uM	12.0% @1.0uM		
B-0146	77.0% @10.0uM	28.0% @1.0uM		
B-0147	44.0% @1.0uM	22.0% @1.0uM		
B-0148	51.0% @1.0uM	>1.0uM		
B-0149	1.15 uM	10.0 uM		
B-0150	27.0% @10.0uM	35.0% @1.0uM		
B-0151	43.0% @1.0uM	30.0% @1.0uM		
B-0152	51.0% @1.0uM	24.0% @1.0uM		
B-0153	57.0% @1.0uM	21.0% @1.0uM		
B-0154	65.0% @10.0uM	14.0% @1.0uM		
B-0155	40.0% @10.0uM	26.0% @1.0uM		
B-0156	42.0% @10.0uM	13.0% @1.0uM		
B-0157	48.0% @10.0uM	9.0% @1.0uM		
B-0158	58.0% @10.0uM	39.0% @1.0uM		
B-0159	54.0% @10.0uM	5.0% @1.0uM		
B-0160	59.0% @10.0uM	26.0% @1.0uM		
B-0161	72.0% @10.0uM	13.0% @1.0uM		
B-0162	23%@1.0uM	2.05 uM		
B-0163	20.0% @10.0uM	10.0% @1.0uM		
B-0164	37.0% @10.0uM	20.0% @1.0uM		
B-0165	70.0% @10.0uM	19.0% @1.0uM		
B-0166	45.0% @10.0uM	37.0% @1.0uM		
B-0167	40.0% @1.0uM	37.0% @1.0uM		
B-0168	44%@1.0uM	2.36 uM		···
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l i	P38 alpha kinase	U937 Ceil IC50.uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	Inhib @dose
	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#				
B-0169	43.0% @1.0uM	21.0% @1.0uM		
B-0170	43.0% @ 1.0uM	30.0% @1.0uM		
B-0171	61.0% @ 10.0uM	21.0% @ 1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0172	16.0% @10.0uM	11.0% @1.0uM		
B-0173	33.0% @10.0uM	48.0% @1.0uM		
B-0174	54.0% @10.0uM	43.0% @1.0uM		
B-0175	41.0% @ 10.0uM	31.0% @1.0uM		
B-0176	50.0% @1.0uM	30.0% @1.0uM		
B-0177	70.0% @10.0uM	27.0% @1.0uM		
B-0178	12.0% @10.0uM	35.0% @1.0uM		
	27.0% @ 10.0uM	37.0% @1.0uM		
B-0180	34.0% @ 10.0uM	23.0% @1.0uM		
B-0181	5.0%@1.0uM	2.0% @1.0uM		
B-0182	39.0% @10.0uM	40.0% @1.0uM		
B-0183	12.0% @10.0uM	34.0% @1.0uM		
B-0184	66.0% @10.0uM	17.0% @1.0uM		
B-0185	65.0% @10.0uM	25.0% @1.0uM		
B-0186	40.0% @ 1.0uM	25.0% @ 1.0uM		
B-0187	4.0% @10.0uM	14.0% @1.0uM		
B-0188	70.0% @ 10.0uM	35.0% @1.0uM		
B-0189	42.0% @10.0uM	9.0% @1.0uM		
B-0190	59.0% @10.0uM	31.0% @1.0uM		
B-0191	40.0% @1.0uM	29.0% @1.0uM		
B-0192	12.0% @10.0uM	47.0% @ 1.0uM		
B-0193	0.54 uM	6%@1.0uM		
B0194	1.31 uM	22%@1.0uM		
B-0195	1.03 uM	55%@1.0uM		
B-0196	2.24 uM	>1.0uM		
B-0197	2.0 uM	14%@1.0uM		
B-0198	1.2 uM	2%@1.0uM		
B-0199	1.34 uM	3%@1.0uM		
B-0200	1.31 uM	16%@1.0uM		
B-0201	0.29 uM	59%@1.0uM		
B-0202	0.55 uM	2.26 uM		
B-0203	0.16 uM	65%@1.0uM		
B-0204	0.21 uM	48%@1.0uM		
B-0205	0.096 uM	54%@1.0uM		
B-0206	5.76 uM	14%@1.0uM		
B-0207	0.12 uM	52%@1.0uM		
B-0208	0.067 uM	>1.0uM		
B-0209	0.29 uM	8%@1.0uM		
B-0210	0.057 uM	67%@1.0uM		

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	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	Inhib @dose
Example#	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
B-0211	0.25 uM	209/ @4 014		
B-0211	0.12 uM	30%@1.0uM		
B-0212 B-0213	0.12 uM	28%@1.0uM		
	0.31 uM	39%@1.0uM		
B-0214		50%@1.0uM		
B-0215	0.11 uM	51%@1.0uM		
B-0216	0.56 uM	>1.0uM		- · · · · · · · · · · · · · · · · · · ·
B-0217	0.55 uM	>1.0uM		
B-0218	0.53 uM	18%@1.0uM		
B-0219	0.91 uM	18%@1.0uM		
B-0220	0.13 uM	40%@1.0uM		
B-0221	2.4 uM	>1.0uM		
B-0222	0.4uM	29.0%@1.0uM		
B-0223	0.2uM	1.0%@1.0uM		
B-0224	<0.1uM	93.0%@1.0uM		
B-0225	0.047uM	37.0%@1.0uM		
B-0226	0.074uM	20.0%@1.0uM		
B-0227	0.045uM	1.0%@1.0uM		
B-0228	. 0.15uM	44.0%@1.0uM		
B-0229	<0.1 uM	61.0%@1.0uM		
B-0230	0.041uM	30.0%@1.0uM		
B-0231	0.055uM	40.0%1.0uM		
B-0232	0.048uM	24.0%@1.0uM		···
B-0233	0.095uM	43.0%@1.0uM		
B-0234	0.11uM	68.0%@1.0uM		
B-0235	1.31uM	90.0%@1.0uM		
B-0236	0.077uM	46.0%@1.0uM		
B-0237	0.13uM	60.0%@1.0uM		*
B-0238	0.47uM	82.0%@1.0uM		
B-0239	5.73uM	84.0%@1.0uM		
B-0240	0.2uM	70.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0241	0.1uM	45.0%@1.0uM		
B-0242	<0.1uM	78.0%@1.0uM		
B-0243	0.039uM	53.0%@1.0uM		
B-0244	0.02uM	57.0%@1.0uM		
B-0245	0.13uM	24.0%@1.0uM		
B-0246	<0.1uM	>1.0uM		
B-0247	0.082uM	75.0%@1.0uM		
B-0248	<0.1uM	11.0%@1.0uM		
B-0249	<0.1uM	75.0%@1.0uM		·
B-0250	0.28uM	36.0%@1.0uM		
B-0251	0.31uM	1.0%@1.0uM		
B-0252	0.041µM	54.0%@1.0uM		
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Example#	P38 aipha kinase iC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or % inhib@conc. (uM)	Mouse LPS Model % TNF Inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-0253	0.061uM	74.0%@1.0uM		
B-0253 B-0254	0.12uM	59.0%@1.0uM		
B-0255	0.12uM	59.0%@1.0uM		
B-0256	<0.1uM			
B-0257	1.71uM	88.0%@1.0uM 11.0%@1.0uM		
B-0258	0.37uM			
B-0258	0.35 uM	63.0%@1.0uM		
B-0259 B-0260	0.55uM 0.56uM	58.0%@1.0uM 23.0%@1.0uM		
B-0260 B-0261	0.49uM			
	0.49uM	23.0%@1.0uM		
B-0262	0.62uM	89.0%@1.0uM		
B-0263 B-0264	0.62uM 0.14uM	64.0%@1.0uM		
B-0265	0.14uM 0.92uM	18.0%@1.0uM		
B-0266	0.92UM 0.25uM	24.0%@1.0uM		
	0.25uM 0.48uM	24.0%@1.0uM		
B-0267	3.39uM	11.0%@1.0uM		
B-0268		19.0%@1.0uM		
B-0269	9.81uM	19.0%@1.0uM		
B-0270	5.79uM	13.0%@1.0uM		
B-0271	7.55uM	12.0%@1.0uM		
B-0272	1.81uM	48.0%@1.0uM		
B-0273	5.03uM	13.0%@1.0uM		
B-0274	2.68uM	25.0%@1.0uM	-	
B-0275	2.67uM	33.0%@1.0uM		
B-0276	1.25uM	26.0%@1.0uM		
B-0277	0.68uM	34.0%@1.0uM		
B-0278	1.26uM	36.0%@1.0uM		
B-0279	1.39uM	33.0%@1.0uM		
B-0280	0.86uM	18.0%@1.0uM		
B-0281	7.37uM	24.0%@1.0uM		
B-0282	0.75uM	38.0%@1.0uM		
B-0283	6.66uM	29.0%@1.0uM		
B-0284	0.083uM	65.0%@1.0uM		
B-0285	4.57uM	29.0%@1.0uM		
B-0286	0.33uM	50.0%@1.0uM		
B-0287	4.0uM	22.0%@1.0uM		
B-0288	4.46uM	26.0%@1.0uM		
B-0289	0.15uM	55.0%@1.0uM		
B-0290	0.66uM	44.0%@1.0uM		
B-0291	1.33uM	20.0%@1.0uM		
B-0292	0.22uM	28.0%@1.0uM		
B-0293	0.66uM	53.0%@1.0uM		
B-0294	0.68uM	45.0%@1.0uM		

in	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM	Mouse LPS Model %	D=41 D0 14- 3.15:
in	IC50.uM or %			Rat LPS Model %
in		or %	TNF Inhib @ dose	Inhib @dose
	hib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#	0.00-11	47.00(04.0.11		
B-0295	0.82uM	45.0%@1.0uM		
B-0296	8.03uM	36.0%@1.0uM		
B-0297	0.78uM	30.0%@1.0uM		
B-0298	0.58uM	48.0%@1.0⊔M		
B-0299	0.87uM	54.0%@1.0uM		
B-0300	0.78uM	32.0%@1.0uM		
B-0301	0.19uM	50.0%@1.0uM		
B-0302	4.02uM	24.0%@1.0uM		
B-0303	0.22uM	10.0%@1.0uM		
B-0304	0.56uM	28.0%@1.0uM		
B-0305				
B-0306				
B-0307				
B-0308				
B-0309				·
B-0310				
B-0311	- 1			
B-0312				
B-0313				
B-0314				
B-0315	<u> </u>			
B-0316				
B-0317				
B-0318				····
B-0319				
B-0320				
B-0321				
B-0322				
B-0323				
B-0324				
B-0325				
B-0326	·			
B-0327				
B-0328				
B-0329				
B-0330				
B-0331				
B-0332				
B-0333	 			
				
B-0334				
B-0335				
B-0336		<u></u>		

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	P38 alpha kinase IC50,uM or %	U937 Ceil IC50,uM or %	Mouse LPS Model % TNF inhib @ dose	Rat LPS Model % inhib @dose
li	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#				
B-0337			ļ	
B-0338			<u> </u>	
B-0339		-,		
B-0340				
B-0341				
B-0342		·		
B-0343				
B-0344				
B-0345				
B-0346				
B-0347			·	
B-0348				
B-0349				
B-0350				
B-0351				
B-0352				
B-0353	1.37uM	55%@1.0uM		
B-0354	1.0uM	0.66uM	51%@30mpk@-6h	54%@3mpk@-4h
B-0355	0.75uM	40.0%@1.0uM		
B-0356	0.66uM	24.0%@1.0uM		
B-0357	1.46uM	0.66uM		
B-0358	0.37uM	17.0%@1.0uM		
B-0359	0.45uM	47.0%@1.0uM		
B-0360	1.6uM	19.0%@1.0uM	·	 .,
B-0361	0.33uM	46.0%@1.0uM		
B-0362	0.52uM	27.0%@1.0uM		
B-0363	4.67uM	25.0%@1.0uM		
B-0364	1.44uM	27.0%@1.0uM		
B-0365	0.96uM	27.0%@1.0uM		
B-0366	0.7uM	46.0%@1.0uM		
B-0367	1.0uM	23.0%@1.0uM		
B-0368	1.0uM	0.64uM	37%@30mpk@-6h	
B-0369	0.16uM	57.0%@1.0uM		
B-0370	0.65uM	28.0%@1.0u M		
B-0371	0.49uM	28.0%@1.0uM		
B-0372	0.35uM	29.0%@1.0uM		·
B-0373	0.45uM	18.0%@1.0uM		
B-0374	1.38uM	12.0%@1.0uM		
B-0375	1.0uM	19.0%@1.0uM		
B-0376	2.99uM	12.0%@1.0uM		
B-0377	1.29uM	36.0%@1.0uM		
B-0378	1.1uM	36.0%@1.0uM		
- 55,6	********	20.0/68 I.OUM	<u></u>	

Example#	P38 alpha kinase IC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or % inhib@conc. (uM)	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-0379	0.53uM	24.0%@1.0uM		
B-0380	1.41uM	32.0%@1.0uM		
B-0381	0.22uM	47.0%@1.0uM		
B-0382	0.41uM	32.0%@1.0uM		
B-0383	1.43uM	10.0%@1.0uM		
B-0384	4.02uM	16.0%@1.0uM		
B-0385	0.057uM	0.9uM	30%@30mpk@-6h	0%@3mpk@-4h
B-0386	0.13uM	54.0%@1.0uM		070001111111111111111111111111111111111
B-0387	0.41uM	52.0%@1.0uM		" ·
B-0388	<0.1uM	36.0%@1.0uM		
B-0389	0.01 uM	0.05uM		62%@3mpk@-4h
B-0390	0.089uM	55.0%@1.0uM		- Carrottiphe III
B-0391	0.86uM	18.0%@1.0uM		
B-0392	0.13uM	57.0%@1.0uM		
B-0393	0.043uM	66.0%@1.0uM		
B-0394	0.13uM	45.0%@1.0uM		·
B-0395	0.087uM	48.0%@1.0uM		
B-0396	0.097uM	0.44uM		
B-0397	0.17uM	41.0%@1.0uM		
B-0398	0.054uM	66.0%@1.0uM		
B-0399	0.14uM	39.0%@1.0uM		
B-0400	0.16uM	25.0%@1.0uM		*****
B-0401	0.46uM	52.0%@1.0uM		
B-0402	0.14uM	1.51uM		
B-0403	1.77uM	2.42uM		
B-0404	0.31uM	48.0%@1.0uM		
B-0405	0.79uM	30.0%@1.0uM		
B-0406	0.54uM	35.0%@1.0uM		
B-0407	0.76uM	27.0%@1.0uM		
B-0408	0.5uM	50.0%@1.0uM		
B-0409	0.53uM	30.0%@1.0uM		
B-0410	0.38uM	44.0%@1.0uM		
B-0411	0.62uM	50.0%@1.0uM		
B-0412	0.24uM	48.0%@1.0uM		
B-0413	0.18uM	55.0%@1.0uM		
B-0414	2.54uM	25.0%@1.0uM		
B-0415	0.42uM	43.0%@1.0uM		
B-0416	0.32uM	34.0%@1.0uM		
B-0417	0.91uM	28.0%@1.0uM		
B-0418	0.22uM	27.0%@1.0uM		
B-0419	0.85uM	41.0%21.0uM		
B-0420	0.83uM	49.0%@1.0uM		

	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM or %	Mouse LPS Model %	
Evample#	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@ predose time
Example# B-0421	0.46uM	57.0%@1.0uM		
B-0421	<0.1uM	40.0%@1.0uM		
B-0422	0.18uM			
B-0423	0.18um	33.0%@1.0uM		
B-0424	0.26uM	32.0%@1.0uM		ļ
B-0425		54.0%@1.0uM		
B-0426	0.055uM	0.74uM		41%@3mpk@-4h
	0.63uM	39.0%@1.0uM		
B-0428	0.99uM	27.0%@1.0uM		
B-0429	0.27uM	45.0%@1.0uM		
B-0430	0.29uM	75.0%@1.0uM		
B-0431	0.21uM	64.0%@1.0uM		
B-0432	<0.1uM	89.0%@1.0uM		
B-0433	<0.1uM	92.0%@1.0uM		
B-0434	0.12uM	65.0%@1.0uM		
B-0435	0.3uM	61.0%@1.0uM		
B-0436	1.11uM	71.0%@1.0uM		
B-0437	0.58uM	59.0%@1.0uM		
B-0438	<0.1uM	91.0%@1.0uM		
B-0439	2.12uM	65.0%@1.0uM		
B-0440	0.66uM	63.0%@1.0uM		
B-0441	0.8uM	58.0%@1.0uM		
B-0442	<0.1uM	91.0%@1.0uM		
B-0443	2.01uM	71.0%@1.0uM		
B-0444	1.01uM	51.0%@1.0uM		
B-0445	<0.1uM	83.0%@1.0uM		
B-0446	0.78uM	80.0%@1.0uM		
B-0447	0.19uM	71.0%@1.0uM		
B-0448	0.4uM	79.0%@1.0uM		
B-0449	0.83uM	81.0%@1.0uM		
B-0450	0.26uM	81.0%@1.0uM		
B-0451	0.071 uM	83.0%@1,0uM	42%@30mpk@-6h	
B-0452	0.7uM	75.0%@1.0uM		
B-0453	0.47uM	75.0%@1.0uM		
B-0454	0.11uM	80.0%@1.0uM		
B-0455	<0.1uM	95.0%@1.0uM		36%@3mpk%-4h
B-0456	1.81uM	67.0%@1.0uM	· ·	30 /0 8 3 HPK /0-413
B-0457	0.089uM	81.0%@1.0uM		
B-0458	0.033uM	70.0%@1.0uM		
B-0459	0.099uM	76.0%@1.0uM		
B-0460	0.061uM	92.0%@1.0uM		
B-0461	0.025uM	96.0%@1.0uM		
B-0462	<0.1uM	97.0%@1.0uM		
- 0702	CO. TURY	31.0%W1.UUM		

Evamuia #	P38 alpha kinase IC50,uM or % Inhib@conc. (uM)	U937 Cell IC50,uM or % inhib@conc. (uM)	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % Inhib @dose @predose time
Example#	0.050-14	22.22.2		
B-0463 B-0464	0.052uM	95.0%@1.0uM		
	<0.1uM	91.0%@1.0uM		
B-0465	0.084uM	98.0%@1.0uM		
B-0466	<0.1uM	98.0%@1.0uM		0%@3mpk@-4h
B-0467	<0.1uM	77.0%@1.0uM		
B-0468	0.031uM	93.0%@1.0uM		
B-0469	0.056uM	92.0%@1.0uM		
B-0470	0.063uM	92.0%@1.0uM		
B-0471	0.027uM	97.0%@1.0uM		
B-0472	0.19uM	54.0%@1.0uM		
B-0473	0.004uM	95.0%@1.0uM		
B-0474	0.024uM	86.0%@1.0uM		
B-0475	0.21uM	74.0%@1.0uM		
B-0476	0.56uM	69.0%@1.0uM		
B-0477	1.48uM	96.0%@1.0uM		
B-0478	0.034uM	87.0%@1.0uM		
B-0479	0.031uM	90.0%@1.0uM		15%@3mpk@-4h
3-0480	0.12uM	Mu0.19%0.88		•
3-0481	0.014uM	95.0%@1.0uM		56%@3mpk@-4h
3-0482	0.97uM	68.0%@1.0uM		
3-0483	0.57uM	68.0%@1.0uM		
3-0484	0.28uM	62.0%@1.0uM		···
3-0485	0.04uM	95.0%@1.0uM		
3-0486	0.24uM	80.0%@1.0uM		
3-0487	0.11uM	89.0%@1.0uM		54%@3mpk@-4h
-0488	0.62uM	88.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
-0489	0.3uM	80.0%@1.0uM		
-0490	0.91uM	74.0%@1.0uM		
-0491	0.43uM	66.0%@1.0uM		
-0492	0.069uM	42.0%@1.0uM		
-0493	0.3uM	36.0%@1.0uM		
-0494	0.13uM	30.0%@1.0uM		···
-0495	0.12uM	25.0%@1.0uM		
-0496	0.83uM	16.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
-0497	0.44uM	31.0%@1.0uM		
-0498	0.33uM	11.0%@1.0uM		
-0499	0.39uM	37.0%@1.0uM		
-0500	0.26uM	41.0%@1.0uM		······································
-0501	0.049uM	52.0%@1.0uM		
-0502	0.065uM	48.0%@1.0uM		
-0503	0.16uM	73.0%@1.0uM		
0504	0.4uM	43.0%@1.0uM		

	P38 alpha kinase	U937 Celi IC50,uM		Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	Inhib @dose
Example#	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
B-0505	0.28uM	44.0%@1.0uM		
B-0506	0.94uM	43.0%@1.0uM		
B-0507	0.18uM			
		75.0%@1.0uM		
B-0508	2.0uM	48.0%@1.0uM		
B-0509	0.1uM	86.0%@1.0uM		
B-0510	0.69uM	61.0%@1.0uM		
B-0511	0.007uM	90.0%@1.0uM		
B-0512	1.0uM	53.0%@1.0uM		
B-0513	0.72uM	52.0%@1.0uM		
B-0514	0.14uM	87.0%@1.0uM		·····
B-0515	0.42uM	61.0%@1.0uM		
B-0516	0.37uM	84.0%@1.0uM		
B-0517	0.094uM	52.0%@1.0uM		
B-0518	0.11uM	64.0%@1.0uM		
B-0519	0.043uM	87.0%@1.0uM		
B-0520	Q.4uM			
B-0521	1.37uM	67.0%@1.0uM		
B-0522		52.0%@1.0uM		
	0.15uM	75.0%@1.0uM		
B-0523	0.19uM	83.0%@1.0uM		
B-0524	0.4uM	77.0%@1.0uM		
B-0525	0.16uM	76.0%@1.0uM		
B-0526	0.031uM	87.0%@1.0uM		-
B-0527	1.09uM	63.0%@1.0uM		
B-0528	0.14uM	70.0%@1.0uM		
B-0529	0.11uM	73.0%@1.0uM		
B-0530	5.53uM	45.0%@1.0uM		
B-0531	0.5uM	48.0%@1.0uM		
B-0532	0.45uM	1.01uM	41%@30mpk@-6h	
B-0533	1.23uM	47.0%@1.0uM		
B-0534 B-0535	0.41uM	54.0%@1.0uM		
B-0535	0.44uM	0.87uM		
B-0536 B-0537	0.46uM 3.44uM	0.15uM		
B-0538	1.13uM	51.0%@1.0uM 45.0%@1.0uM		
B-0539	2.84uM	21.0%@1.0uM	· · · · · · · · · · · · · · · · · · ·	
B-0540	3.62uM	54.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0541	3.24uM	28.0%@1.0uM		
B-0542	1.55uM	50.0%@1.0uM		
B-0543	1.56uM	43.0%@1.0uM		
B-0544	1.12uM	27.0%@1.0uM		
B-0545 B-0546	1.06uM	41.0%@1.0uM		
B-0547	1.04uM	18.0%@1.0uM		
B-0548	1.24uM 1.77uM	21.0%@1.0uM		
B-0549	2.22uM	28.0%@1.0uM 22.0%@1.0uM		
	4.EEUIVI	22.0 /6 W 1.UUM		

	P38 alpha kinase IC50.uM or %	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	inhib@conc. (uM)	Inhib@conc. (uM)	Opredose time	inhib @dose @predose time
Example#	mina e conc. (am)	mino e conc. (am)	e predose time	e predose time
B-0550	2.41uM	14.0%@1.0uM		
B-0551	1.08uM	56.0%@1.0uM		
B-0552	0.13uM	46.0%@1.0uM		
B-0553	1.44uM	47.0%@1.0uM		
B-0554	2.58uM	20.0%@1,0uM	-	
B-0555	1.87uM	34.0%@1.0uM		
B-0556	0.49uM	39.0%@1.0uM		
B-0557	1.37uM	32.0%@1.0uM		
B-0558	0.85uM	33.0%@1.0uM		
B-0559	0.53uM	49.0%@1.0uM		· · · · ·
B-0560	2.57uM	31.0%@1.0uM		
3-0561	2.07uM	40.0%@1.0uM		
3-0562	0.22uM	0.3uM		5%@3mpk@-4h
3-0563	0.18uM	0.13uM		
B-0564	0.82uM	58%@1.0uM		
B-0565	0.23uM	0.59uM		
B-0566	<0.1uM	0.17uM		0%@3mpk@-4
B-0567	0.14uM	0.28uM		C/CCCIII.pite 4
B-0568	1.22uM	46.0%@1.0uM		
3-0569	0.15uM	0.26uM		
3-0570	0.27uM	46.0%@1.0uM		
3-0571	0.38uM	44.0%@1.0uM		
3-0572	0.27uM	41.0%@1.0uM		
B-0573	0.36uM	1.7uM		
B-0574	0.13uM	0.66uM		37%@3mpk@-4
B-0575	0.032uM	0.17u M		
3-0576	0.068uM	0.39uM		65%@3mpk@-4
B-0577	0.091uM	66.0%@1.0uM		CO / C C C III PICC 1
3-0578	1.88uM	47.0%@1.0uM		
3-0579	0.11uM	79.0%@1.0uM		
3-0580	2.23uM	0.84uM		
3-0581	0.26uM	2.17uM		
3-0582	1.03uM	37.0%@1.0uM		
3-0583	3.93uM	26.0%@1.0uM		
3-0584	0.66uM	54.0%@1.0uM		
3-0585	0.83uM	79.0%@1.0uM	50%@30mpk@-6h	
3-0586	0.81uM	51.0%@1.0uM		
3-0587	6.84uM	38%@1.0uM		
3-0588	12.8uM	42%@1.0uM		
3-0589	1.71uM	42%@1.0uM		
3-0590	1.57uM	38.0uM		
3-0591	3.59uM	29.0%@1.0uM		
3-0592	1.62uM	45.0%@1.0uM		
3-0593	1.22uM	36.0%@1.0uM		
3-0594		41.0%@1.0uM		
3-0595	2.42uM	22.0%@1.0uM		
3-0596	20.0uM	41.0%@1.0uM		
3-0597	1.68uM	63.0%@1.0uM		
3-0598	2.12uM	50.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·

r				
1	P38 alpha kinase	U937 Cell 1C50.uM	Mouse LPS Model %	Rat LPS Modei %
1 1	IC50,uM or %	or %	TNF inhib @ dose	inhib @dose
	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#	(4.1.)		o produce timo	o predose time
B-0599	4.16uM	21.0%@1.0uM		
B-0600	0.002 uM	28.0%@1.0uM		
B-0601	0.089uM	1.31uM		43%@3mpk%-4h
B-0602	0.97uM	61.0%@1.0uM		
B-0603	0.09uM	51.0%@1.0uM		
B-0604	0.3uM	20.0%@1.0uM		
B-0605	0.18uM	47.0%@1.0uM		
B-0606	0.17uM	53.0%@1.0uM		
B-0607	2.79uM	70.0%@1.0uM		
B-0608	0.059uM	73.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0609	<0.1uM	87.0%@1.0uM		
B-0610	<0.1uM	88.0%@1.0uM		
B-0611	0.65uM	60.0%@1.0uM		
B-0612	0.16uM	60.0%@1.0uM		
B-0613	0.17uM	76.0%@1.0uM		
B-0614	0.76uM	70.0%@1.0uM		0%@3mpk@-4h
B-0615	0.08uM	83.0%@1.0uM		0 Wednipke 411
B-0616	0.38uM	87.0%@1.0uM		
B-0617	0.045uM	92.0%@1.0uM		
B-0618	0.37uM	80.0%@1,0uM		
B-0619	<0.1uM	88.0%@1.0uM		
B-0620	1.59uM	58.0%@1.0uM		
B-0621	0.36uM	68.0%@1.0uM		
B-0622	0.076uM	78.0%@1.0uM		
B-0623	0.12uM	76.0%@1.0uM		····
B-0624	0.085uM	54.0%@1.0uM	· · · · · · · · · · · · · · · · · · ·	
B-0625	0.023uM	88.0%@1.0uM		
B-0626	<0.1 uM	85.0% @ 1.0uM		
B-0627	0.25uM	69.0%@1.0uM		
B-0628	0.023uM	72.0%@1.0uM		
B-0629	0.2uM	79.0%@1.0uM		
B-0630	0.06uM	77.0%@1.0uM	·	
B-0631	0.065uM	81.0%@1.0uM		
B-0632	<0.1uM	79.0%@1.0uM		
B-0633	0.6uM	80.0%@1.0uM		
B-0634	0.6uM	40.0%@1.0uM		
B-0635	0.15uM	55.0%@1.0uM		
B-0636	<0.1uM	86.0%@1.0uM		
B-0637	0.11uM	92.0%@1.0uM		······································
B-0638	0.25uM	89.0%@1.0uM		
B-0639	0.051 uM	93.0%@1.0uM		50%@3mpk@-4h
B-0540	0.36uM	94.0%@1.0uM		
B-0641	0.58uM	65.0%@1.0uM		
B-0642	0.49uM	90.0%@1.0uM		
B-0643	0.069uM	85.0%@1.0uM		0%@3mpk@-4h
B-0644	0.058uM	89.0%@1.0uM		F
B-0645	0.58uM	80.0%@1.0uM		
B-0646				
B-0647	0.26uM	94.0%@1.0uM		

	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	Inhib @dose
	Inhib@conc. (uM)	inhib@conc. (uM)	© predose time	
Example#	minute conc. (dis)	innibecone. (um)	e predose time	@predose time
B-0648	<0.1uM	83.0%@1.0uM		
B-0649	0.83uM	39.0%@1.0uM		
B-0650	0.006uM	95.0%@1.0uM		8%@3mpk@-4h
B-0651	1.78uM	81.0%@1.0uM		O/GOOMPRO TH
B-0652	0.19uM	83.0%@1.0uM		
B-0653	2.01uM	74.0%@1.0uM		
B-0654	5.97uM	78.0%@1.0uM		
B-0655	1.25uM	76.0%@1.0uM		
B-0656	0.007uM	95.0%@1.0uM		28%@3mpk@-4h
B-0657	0.17uM	83.0%@1.0uM		action of the state of the stat
B-0658	1.14uM	91.0%@1.0uM		
B-0659	2.64uM	87.0%@1.0uM		
B-0660	0.088uM	92.0%@1.0uM		·
B-0661	<0.1uM	90.0%@1.0uM		
B-0662	<0.1uM	95.0%@1.0uM		
B-0663	0.88uM	74.0%@1.0uM		
B-0664	0.39uM_	80.0%@1.0uM		
B-0665	0.47uM	72.0%@1.0uM		
B-0666	0.17uM	73.0%@1.0uM		
B-0667	0.83uM	75.0%@1.0uM		
B-0668	0.27uM	78.0%@1.0uM		
B-0669	0.89uM	34.0%@1.0uM		
B-0670	3.15uM	32.0%@1.0uM		
B-0671	6.38uM	36.0%@1.0uM		
8-0672	6.59uM	32.0%@1.0uM		
B-0673	8.54uM	48.0%@1.0uM		
B-0674	2.81uM	42.0%@1.0uM		
B-0675	5.42uM	3.0%@1.0uM		
B-0676	2.09uM	22.0%@1.0uM		
B-0677	1.63uM	25.0%@1.0uM		
B-0678	0.38uM	52.0%@1.0uM		
B-0679 B-0680	0.062uM	45.0%@1.0uM		
B-0681	0.42uM	67.0%@1.0uM		
B-0682	1.96uM	17.0%@1.0uM		
B-0683	0.76uM	39.0%@1.0uM		
B-0684	13.0uM	32.0%@1.0uM		
3-0685	0.54uM 15.4uM	68.0%@1.0uM		
3-0686		33.0%@1.0uM		
3-0687	0.42uM 10.1uM	59.0%@1.0uM		
3-0688	0.66uM	15.0%@1.0uM		
3-0689		58.0%@1.0uM		
3-0690	14.6uM 27.1uM	27.0%@1.0uM		
3-0691	0.16uM	36.0%@1.0uM		
3-0692	0.18uM	48.0%@1.0uM		
3-0693	0.39uM	29.0%@1.0uM		
3-0694	0.62uM	28.0%@1.0uM		
3-0695	0.82UM 0.23uM	21.0%@1.0uM		
3-0696	0.085uM	32.0%@1.0uM		
	U.UGDU.U	35.0%@1.0uM		

	P38 aipha kinase IC50,uM or %	U937 Cell (C50,uM	Mouse LPS Model %	Rat LPS Model % inhib @dose
	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#			•	
B-0697	0.45uM	44.0%@1.0uM		
B-0698	2.33uM	43.0%@1.0uM		
B-0699	0.34uM	31.0%@1.0uM		·
B-0700	0.24uM	56.0%@1.0uM		
B-0701	0.39uM	45.0%@1.0uM		
B-0702	0.036uM	39.0%@1.0uM		···
B-0703	0.12uM	39.0%@1.0uM		
B-0704	2.19uM	29.0%@1.0uM		
B-0705	0.44uM	21.0%@1.0uM		
B-0706	0.44uM	32.0%@1.0uM		
B-0707	1.7uM			
B-0708	2.1uM			
B-0709	0.84uM			
B-0710	1.99uM			
B-0711	1,99uM			
B-0712	2.9uM			
B-0713	4.3uM			
B-0714	3.7uM			
B-0715	3.2uM			
B-0716	4.6uM			
B-0717	4.3u M			
B-0718	1.4uM			
B-0719	3.4uM			
B-0720	1.3uM			
B-0721	3.8uM			
B-0722	0.07uM	>1.0uM		
B-0723	0.47uM			
B-0724	0.06uM	17.0%@1.0uM		
3-0725	9.7uM			
3-0726	1.4uM			
3-0727	0.51uM			
3-0728	20.0uM			
3-0729	0.87uM			
3-0730	0.25uM	11.0%@1.0uM		
3-0731	0.87uM	>1.0uM		
3-0732	14.0uM			
3-0733	32.0uM			
3-0734	0.92uM			
3-0735	1.0uM			
3-0736	26.0uM			
3-0737	2.6uM			
3-0738	2.7uM			
3-0739	4.1 uM			
3-0740	4.4uM		· · · · · · · · · · · · · · · · · · ·	
3-0741	26.0uM			
3-0742	2.2uM			
3-0743	1,2uM			
3-0744	23.0uM	·		
-0745	6.0uM			

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	P38 alpha kinase	U937 Cell 1C50,uM	Mouse LPS Model %	Rat LPS Model %
i I	IC50,uM or %	or %	TNF inhlb@dose	inhib @dose
	Inhib@conc. (uM)	inhib@conc. (uM)	@predose time	epredose time
Example#	minus econor (um)	minbecone. (um)	e bredose time	e predose time
B-0746	0.01uM	22.0%@1.0uM		
B-0747	1.1uM	Later to Criticalii		
B-0748	1.2uM			
B-0749	4.4uM			
B-0750	0.92uM			
B-0751	1.6uM			
B-0752	0.33uM			
B-0753	0.37uM			· · · · · · · · · · · · · · · · · · ·
B-0754	0.55uM			
B-0755	2.3uM			
B-0756	0.94uM			
B-0757	0.54uM	16.0%@1.0µM		
B-0758	1.5uM			
B-0759	0.3uM			·
B-0760	0.01uM	13.0%@1.0uM		
B-0761	<0.1uM			
B-0762	0.13uM	5.0%@1.0uM		
B-0763	0.015uM	17.0%@1.0uM		
B-0764	0.67uM	26.0%@1.0uM		
B-0765	0.3uM	29.0%@1.0uM		
B-0766	0.95uM			
B-0767	Mu80.0			
B-0768	1.4uM			
B-0769	12.7uM			
B-0770	2.3uM			
B-0771	0.5uM			
B-0772	0.8uM			
B-0773	14.0uM			
B-0774	1.5uM			
B-0775	0.6uM	>1.0uM		
B-0776	0.9uM	>1.0uM		
B-0777	21.0uM			
B-0778	51.0uM			
B-0779 B-0780	0.5uM			
	1.1uM			
B-0781 B-0782	48.0uM			
B-0782 B-0783	22.0uM			
B-0784	8.0uM			
B-0785	7.0uM			
B-0786	23.0uM			
B-0787	24.0uM			
B-0788	1.5uM 1.2uM			
B-0789				
B-0790	33.0uM	4.00/ 634.0-14		· · · · · · · · · · · · · · · · · · ·
B-0791	1.0uM	4.0%@1.0uM		
B-0792	0.3uM 1.1uM	>1.0uM		
B-0793	0.3uM		···	
B-0794	2.9uM	2.09/@4.0-14		
0-0134	Z.9UM	2.0%@1.0uM		

	D20 alpha biass	HOST CON IOTO	Moune I Do Mardal Av	Det I DC Made In
	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM or %	Mouse LPS Model % TNF Inhib @ dose	Rat LPS Model % Inhib @dose
	inhib@conc. (uM)		@predose time	@predose time
Example#				- p
B-0795	1.9uM	11.0%@1.0uM		
B-0796	1.4uM			
B-0797	1.04uM	-		***************************************
B-0798	1.73uM	-		
B-0799		>1.0uM		
B-0800	1.01uM	>1.0uM		
B-0801	0.67uM	>1.0uM		
B-0802	•	>1.0uM		
B-0803	0.057uM	53.0%@1.0uM		
B-0804	0.3uM	32.0%@1.0uM		
B-0805	0.71uM	>1.0uM		
B-0806	3.28uM	>1.0uM		
B-0807	10.8uM	•		
B-0808	3.09uM	>1.0uM		
B-0809	1.22uM	7.0%@1.0uM		
B-0810	1.11uM	>1.0uM		
B-0811	2.79uM	2.0%@1.0uM		
B-0812	2.12uM	>1.0uM		
B-0813	3.02uM	>1.0uM		
B-0814	•	>1.0uM		
B-0815	2.11uM	>1.0uM		
B-0816	3,46uM	>1.0uM		
B-0817	3.07uM	33.0%@1.0uM		
B-0818	4.97uM	>1.0uM		
B-0819	1.08uM	>1.0uM		
B-0820	1.64uM	3.0%@1.0uM		
B-0821	1.44uM			<u> </u>
B-0822	1.33uM			
B-0823	2.39uM	>1.0uM		
B-0824	3.41uM	•		
B-0825		<u> </u>		
B-0826	1.74uM	•		
B-0827	15.6uM	-		
B-0828 B-0829	7.9uM	-		
B-0829 B-0830	0.61uM	65.0%@1.0uM		
B-0830 B-0831	0.54uM	34.0%@1.0uM		
B-0832	0.9uM	>1.0uM		
B-0832 B-0833	1.49uM	00.00/.04.0.14		
B-0834	0.95uM	23.0%@1.0uM		
B-0835	1.25uM	•		
B-0836	1.24uM	•		
B-0837	1.24um 1.96uM	- >1.0uM		
B-0838	3.1uM	> 1.00M		
B-0839	4.3uM			
B-0840	0.63uM	47.0%@1.0uM		
B-0841	0.32uM	36.0%@1.0uM		
B-0842	0.32uM 0.74uM	63.0%@1.0uM		
B-0842	0.61uM	>1.0uM		
2 0043	U.O LUIVI 1	>1.UUM	<u> </u>	

	P38 alpha kinase IC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or % Inhib@conc. (uM)	Mouse LPS Model % TNF Inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
Example#				
B-0844	0.4uM	25.0%@1.0uM		
B-0845	1.78uM			
B-0846	1.8uM	-		
B-0847	0.73uM	21.0%@1.0uM		
B-0848	1.56uM	•		
B-0849	1.25uM	•		
B-0850	1.81uM	-		
B-0851	0.91uM	39.0%@1.0uM		
B-0852	1.02uM			
B-0853	•	38.0%@1.0uM		
B-0854	•	25.0%@1.0uM		
B-0855	•	8.0%@1.0uM		
B-0856	•	38.0%@1.0uM		
B-0857	6.25uM	•		
B-0858	2,1uM	48.0%@1.0uM		
B-0859	39.5uM	•		
B-0860	38.1uM	*		
B-0861	1.32uM	12,0%@1,0uM		
B-0862	2,15uM	4.0%@1.0uM		
B-0863	0.81uM	25.0%@1.0uM		
B-0864	0,39uM	40,%@1,0uM		
B-0865	0.66uM	46.0%@1.0uM		
B-0866	1.38uM	28.0%@1.0uM		
B-0867	0.62uM	>1.0uM		
B-0868	3.28uM	8.0%@1.0uM		
B-0869	4.19uM	>1.0uM		
8-0870	3.13uM	>1.0uM		
B-0871	1.9uM	>1.0uM		
B-0872	3.13uM	3.0%@1.0uM		
B-0873	6.92uM	>1.0uM		
B-0874	1.92uM	>1.0uM		
B-0875	2.13uM	8%@1.0uM		
B-0876	0.89uM	>1.0uM	· · · · · · · · · · · · · · · · · · ·	
B-0877	1.17uM	13.0%@1.0uM	· · · · · · · · · · · · · · · · · · ·	
B-0878	0.65uM	19.0%@1.0uM	······	
B-0879	0.87uM	1.0%@1.0uM		
B-0880	0.15uM	40.0%@1.0uM	i	
B-0881	1.36uM	>1.0uM		
B-0882	1.48uM	9%@1.0uM		
B-0883	1.06uM	>1.0uM		
B-0884	1.89uM	71100111		
B-0885			· · · · · · · · · · · · · · · · · · ·	
B-0886				
B-0887				
B-0888				
B-0889		· · · · · · · · · · · · · · · · · · ·		
B-0890			l ————————————————————————————————————	
B-0891				
B-0892				

	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	inhib @dose
	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#				
B-0893				
B-0894				
B-0895				
B-0896				
B-0897				
B-0898				
B-0899				
B-0900				
B-0901				
B-0902				
B-0903				
B-0904				····
B-0905				
B-0906				
B-0907				
B-0908				
B-0909	• • • • • • • • • • • • • • • • • • • •			
B-0910		-		
B-0911				
B-0912				
B-0913				
B-0914				
B-0915				
B-0916				
B-0917				
B-0918				
B-0919				
B-0920				
B-0921				
B-0922				
B-0923				
B-0924				
B-0925				
B-0926				
B-0927				
B-0928				
B-0929				
B-0930				
B-0931				
B-0932			ii	
B-0933	47.0%@1.0uM	37.0%@1.0uM		
B-0934	67.0%@1.0uM	36.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0935	69.0%@1.0uM	54.0%@1.0uM		
B-0936	69.0%@1.0uM	>1.0uM		· , · · · · · · · · · · · · · · · · · ·
B-0937	64.0%@1.0uM	1.74uM		
B-0938	51.0%@1.0uM	29.0%@1.0uM	-	
B-0939	78.0%@1.0uM	14.0%@1.0uM		
B-0940	56.0%@1.0uM	22.0%@1.0uM		
B-0941				

f				
	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM or %	Mouse LPS Model % TNF inhib @ dose	Rat LPS Model % inhib @dose
F10#	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#	00.00/ @1.0-4/	0.00/ @4.0034		
B-0942	82.0%@1.0uM	2.0%@1.0uM		
B-0943 B-0944	63.0% @10.0uM	24.0%@1.0uM 27.0%@1.0uM		
B-0944	45.0%@1.0uM	0.93uM		
	96.0%@1.0uM			
B-0946 B-0947	76.0%@1.0uM	31.0%@1.0uM		
	69.0%@1.0uM	34.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-0948 B-0949	68.0%@1.0uM	1.81uM 17.0%@1.0uM		
	90.0%@1.0uM			
B-0950	81.0%@1.0uM	0.58uM		
B-0951 B-0952	82.0%@1.0uM	20.0%@1.0uM		
	44.0%@1.0uM	21.0%@1.0uM		
B-0953	63.0%@1.0uM	25.0%@1.0uM		
B-0954	62.0%@1.0uM	0.52uM		
B-0955	49.0%@1.0uM	0.54uM		
B-0956	56.0%@1.0uM	1.33uM		
B-0957	79.0%@1.0uM	22.0%@1.0uM		
B-0958	74.0%@1.0uM	0.38uM		
B-0959	83.0%@1.0uM	39.0%@1.0uM		
B-0960	48.0%@1.0uM	4.0%@1.0uM		
B-0961	79.0%@1.0uM	23.0%@1.0uM		
B-0962	85.0%@1.0uM	2.71uM		
B-0963	76.0%@1.0uM	39.0%@1.0uM		
B-0964	94.0%@1.0uM	5.0uM		
B-0965	74.0%@1.0uM	1.1uM		
B-0966	50.0%@1.0uM	5.0%@1.0uM		
B-0967	80.0%@1.0uM	29.0%@1.0uM		
B-0968	35.0%@1.0uM	26.0%@1.0uM		
B-0969	63.0%@1.0uM	35.0%@1.0uM		
B-0970	76.0%@10.0uM	0.88uM		
B-0971	61.0%@1.0uM	39.0%@1.0uM		
B-0972	85.0%@1.0uM	2.0%@1.0uM		
B-0973	66.0%@10.0uM	48.0%@1.0uM		
B-0974	57.0%@1.0uM	47.0%@1.0uM		
B-0975	82.0%@1.0uM	32.0%@1.0uM		
B-0976	79.0%@1.0uM	36.0%@1.0uM		
B-0977	60.0%@1.0uM	26.0%@1.0uM		
B-0978	59.0%@1.0uM	36.0%@1.0uM		
B-0979	56.0%@10.0uM	23.0%@1.0uM		
B-0980	68.0%@1.0uM	31.0%@1.0uM		
B-0981	62.0%@1.0uM	57.0%@1.0uM		
B-0982	65.0%@1.0uM	23.0%@1.0uM		
B-0983	75.0%@1.0uM	0.8uM		
B-0984 B-0985	60.0%@1.0uM	51.0%@1.0uM		
B-0986	86.0%@1.0uM	0.75uM		
B-0987	70.0%@1.0uM	71.0%@1.0uM		
B-0988	78.0%@1.0uM 72.0%@1.0uM	79.0%@1.0uM		
B-0989	72.0%@1.0UM 85.0%@1.0UM	65.0%@1.0uM		
B-0989	MDU. W 67U.co	0.85uM		
10-0990	•	26.0%@1.0uM		<u> </u>

ICSO HM or 9/	U937 Cell IC50,uM	Mouse LPS Model % TNF inhib @ dose	Rat LPS Model %
IC50,uM or % inhib@conc. (uM)	inhib@conc. (uM)	1	Inhib @dose
minu & conc. (dwi)	minbeconc. (um)	epredose time	@predose time
58.0%@1.0uM	33.0%@1.0uM		
57.0%@1.0uM			
55.0%@1.0uM			
53.0%@1.0uM			
54.0%@1.0uM	27.0%@1.0uM		
69.0%@1.0uM	22.0%@1.0uM		
67.0%@1.0uM	25.0%@1.0uM		
61.0%@1.0uM	24.0%@1.0uM		
55.0%@1.0uM	42.0%@1.0uM		
63.0%@1.0uM	31.0%@1.0uM		
70.0%@1.0uM	41.0%@1.0uM		
74.0%@1.0uM	29.0%@1.0uM		
79.0%@1.0uM	45.0%@1.0uM		
58.0%@1.0uM	23.0%@1,0uM		*
69.0%@1.0uM	38.0%@1,0uM		
	34.0%@1.0⊔M		
	23.0%@1.0uM		
	55.0%@1.0uM		
	1.0uM		
72.0%21.0uM			
		····	
			·
00.076@1.0UM			
57.0%@1.0044			
- 31.0 /0 @ 1.00M			
	58.0%@1.0uM 77.0%@1.0uM 57.0%@1.0uM 55.0%@1.0uM 53.0%@1.0uM 54.0%@1.0uM 69.0%@1.0uM 67.0%@1.0uM 61.0%@1.0uM 65.0%@1.0uM 70.0%@1.0uM 74.0%@1.0uM 74.0%@1.0uM 78.0%@1.0uM	58.0%@1.0uM	58.0%@1.0uM

	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
Example#	Inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
B-1040	70 00/ 0 0 0 0			
	72.0%@1.0uM	0.38uM		
B-1041	70.0%@1.0uM	73.0%@1.0uM		
B-1042	79.0%@1.0uM	12.0%@1.0uM		
B-1043	64.0%@1.0uM	53.0%@1.0uM		
B-1044	94.0%@1.0uM	0.93uM		
B-1045	78.0%@1.0uM	25.0%@1.0uM		
B-1046	72.0%@1.0uM	66.0%@1.0uM		·
B-1047	72.0%@1.0uM	58.0%@1.0uM		
B-1048	67.0%@1.0uM	19.0%@1.0uM		
B-1049	67.0%@1.0uM	65.0%@1.0uM		
B-1050	•	0.54uM		
B-1051	68.0%@1.0uM	41%@1.0uM		··
B-1052	69.0%@1.0uM	66%@1.0uM		
B-1053	78.0%@1.0uM	0.4uM		
B-1054	79.0%@1.0uM	55.0%@1.0uM		
B-1055	89.0%@1.0uM	63.0%@1.0uM		
B-1056	89.0%@1.0uM	0.76uM		
B-1057	85.0%@1.0uM	0.72uM		
B-1058	0.66uM	43.0%@1.0uM		
B-1059	0.18uM	24.0%@1.0uM		
3-1060	0.11uM	32.0%@1.0uM		
3-1061	0.03uM	19.0%@1.0uM		
3-1062	<0.1uM	26.0%@1.0uM		
3-1063	0.16uM	44.0%@1.0uM		
3-1064	0.39uM	50.0%@1.0uM		
3-1065	0.56uM	40.0%@1.0uM		·
3-1066	<0.1uM	39.0%@1.0uM		
3-1067	1.6uM	32.0%@1.0uM		
3-1068	0.48uM	24.0%@1.0uM		
-1069	0.22uM			
3-1070	<0.1uM	27.0%@1.0uM		
-1071	<0.1uM	44.0%@1.0uM		
-1072	0.38uM	48.0%@1.0uM		
-1073	<0.1uM	28.0%@1.0uM		
-1074		21.0%@1.0uM		
-1075	0.23uM	33.0%@1.0uM		
-1076	0.03uM	29.0%@1.0uM		
-1077	0.08uM	31.0%@1.0uM		
-1078	<0.1uM	38.0%@1.0uM		
-1079	0.26uM	48.0%@1.0uM		
-1079	<0.1uM	40.0%@1.0uM		
-1080	0.19uM	28.0%@1.0uM		
	<0.1uM	37.0%@1.0uM		
1082	<0.1uM	54.0%@1.0uM		
1083	<0.1uM	23.0%@1.0uM		
1084	0.43uM	29.0%@1.0uM		
1085	<0.1uM	29.0%@1.0uM		
1086	<0.1uM	42.0%@1.0uM		
1087	0.05uM	32.0%@1.0uM		
1088	0.73uM	49.0%@1.0uM		

Example# B-1089	use LPS Model % Inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-1090 <0.1 uM 90.0%@1.0 uM B-1091 <0.1 uM 73.0%@1.0 uM B-1092 0.27 uM 85.0%@1.0 uM B-1093 0.33 uM 36.0%@1.0 uM B-1094 0.013 uM 69.0%@1.0 uM B-1095 <0.1 uM 70.0%@1.0 uM B-1096 <0.1 uM 32.0%@1.0 uM B-1096 <0.1 uM 32.0%@1.0 uM B-1097 <0.1 uM 44.0%@1.0 uM B-1098 <0.1 uM 82.0%@1.0 uM B-1099 0.26 uM 74.0%@1.0 uM B-1099 0.26 uM 74.0 @1.0 uM B-1100 0.22 uM 56.0 @1.0 uM B-1101 0.025 uM 82.0 @1.0 uM B-1102 0.035 uM 83.0 %@1.0 uM B-1103 0.094 uM 90.0 %@1.0 uM B-1104 0.12 uM 69.0 %@1.0 uM B-1105 <0.1 uM 84.0 %@1.0 uM B-1106 <0.1 uM 84.0 %@1.0 uM B-1108 0.22 uM 81.0 %@1.0 uM B-1109 0.057 uM 84.0 %@1.0 uM B-1109 0.057 uM 84.0 %@1.0 uM B-1109 0.054 uM 80.0 %@1.0 uM B-1111 0.19 uM 64.0 %@1.0 uM B-1112 0.58 uM 81.0 %@1.0 uM B-1111 0.19 uM 64.0 %@1.0 uM B-1112 0.58 uM 81.0 %@1.0 uM B-1113 0.07 uM 81.0 %@1.0 uM B-1123 0.58 uM 29.0 %@1.0 uM B-1124 1.49 uM 52.0 %@1.0 uM B-1125 0.56 uM 41.0 uM 81.0 uM		
B-1091 <0.1uM 73.0%@1.0uM		
B-1092		
B-1093 0.33uM 36.0%@1.0uM		
B-1094		
B-1095		
B-1096 <0.1uM 32.0%@1.0uM		
B-1097		
B-1097		
B-1099		
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B-1126 1.5uM >1.0uM B-1127 0.71uM 7.0%@1.0uM B-1128 2.55uM 26.0%@1.0uM B-1129 1.07uM 46.0%@1.0uM B-1130 0.5uM 29.0%@1.0uM B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1127 0.71uM 7.0%@1.0uM B-1128 2.55uM 26.0%@1.0uM B-1129 1.07uM 46.0%@1.0uM B-1130 0.5uM 29.0%@1.0uM B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1128 2.55uM 26.0%@1.0uM B-1129 1.07uM 46.0%@1.0uM B-1130 0.5uM 29.0%@1.0uM B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1129 1.07uM 46.0%@1.0uM B-1130 0.5uM 29.0%@1.0uM B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1130 0.5uM 29.0%@1.0uM B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1131 0.076uM 34.0%@1.0uM B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		·
B-1132 0.72uM 11.0%@1.0uM B-1133 0.38uM 33.0%@1.0uM		
B-1133 0.38uM 33.0%@1.0uM		·····
		
		
B-1136 1.17uM 40.0%@1.0uM B-1137 0.038uM 35.0%@1.0uM		

	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
1	IC50,uM or %	or %	TNF inhib @ dose	inhib @dose
	Inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#			o produce time	e predose time
B-1138	1.82uM	>1.0uM		
B-1139	0.041 uM	29.0%@1.0uM		
B-1140	1.68uM	39.0%@1.0uM		
B-1141	2.47uM	32.0%@1.0uM		
B-1142	0.11uM	37.0%@1.0uM		
B-1143	0.17uM	40.0%@1.0uM		
B-1144	0.44uM	72.0%@1.0uM		
B-1145	1.07uM	71.0%@1.0uM		
B-1146	0.47uM	61.0%@1.0uM		
B-1147	0.095uM	53.0%@1.0uM		
B-1148	0.43uM	61.0%@1.0uM		
B-1149	1.55uM	48.0%@1.0uM		
B-1150	0.47uM	75.0%@1.0uM		
B-1151	0.32uM	72.0%@1.0uM		
B-1152	0.73uM	53.0%@1.0uM		
B-1153	2.22uM	52.0%@1.0uM		
B-1154	0.085uM	46.0%@1.0uM		
B-1155	3,22uM	30.0%@1.0uM		
B-1156	0.27uM	78.0%@1.0uM		
B-1157	0.26uM	66.0%@1.0uM		
B-1158	74%@1.0uM	0.68uM	53%@30mpk@-6h	
B-1159	66.0%@1.0uM	1.03uM	60%@30mpk@-6h	
B-1160	79.0%@1.0uM	0.38uM		
B-1161	64.0%21.0uM	0.93uM	40%@30mpk@-6h	45%@3mpk@-4h
B-1162	79.0%@1.0uM	0.59uM	40%@30mpk@-6h	TO TO COMPAGE 411
B-1163	74.0%@1.0uM	0.37uM		*
B-1164		0.35uM		
B-1165	66.0%@1.0uM	0.99uM		
B-1166	77.0%@1.0uM	0.39uM	50%@30mpk@-6h	50%@3mpk@-4h
B-1167	70.0%@1.0uM	1.06uM		
B-1168	66.0%@1.0uM	0.63uM		
B-1169	80.0%@1.0uM	0.11uM		
B-1170	82.0%@1.0uM	0.57uM		
B-1171	78.0%@1.0uM	0.23uM		
B-1172	68.0%@1.0uM	1.95uM		***************************************
B-1173	65.0%@1.0uM	62%@1.0uM		
B-1174	80.0%@1.0uM	0.86uM		
B-1175	72.0%@1.0uM	1.83uM		
B-1176	67.0%@1.0uM	67.0%@1.0uM		
B-1177	70.0%@1.0uM	1.16uM		
B-1178	92.0%@1.0uM	1.61uM		
B-1179	86.0%@1.0uM	0.41uM		
B-1180	78.0%@1.0uM	0.53uM		
B-1181	79.0%@1.0uM	66%@1.0uM		
B-1182	72.0%@1.0uM	0.65uM		
B-1183	77.0%@1.0uM	0.2uM		
B-1184	69.0%@1,0uM	0.63uM		
B-1185	71.0%@1.0uM	0.79uM		
B-1186	83.0%@1.0uM	60%@1.0uM		
-				

	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM or %	Mouse LPS Model %	Rat LPS Model % inhlb @dose
	inhlb@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#				
B-1187	76.0%@1.0uM	1.89uM		
B-1188	•	36.0%@1.0uM		
B-1189	68.0%@1.0uM	0.83uM		
B-1190	78.0%@1.0uM	62.0%@1.0uM		
B-1191	74.0%@1.0uM	57.0%@1.0uM		
B-1192	84.0%@1.0uM	0.47uM		
B-1193	69.0%@1.0uM	65.0%@1.0uM		
B-1194	87.0%@1.0uM	0.58uM		
B-1195	52.0%@1.0uM	60.0%@1.0uM		
B-1196	74.0%@1.0uM	68.0%@1.0uM		
B-1197	77.0%@1.0uM	45.0%@1.0uM		
B-1198	92.0%@1.0uM	0.46uM		
B-1199	87.0%@1.0uM	49.0%@1.0uM		
B-1200	95.0%@1.0uM	0.64uM		
B-1201	84.0%@1.0uM	0.51uM		
B-1202	71.0%@1.0uM	58.0%@1.0uM		
B-1203	84.0%@1.0uM	58.0%@1.0uM		
B-1204	68.0%@1.0uM	59.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-1205	74.0%@1.0uM	46.0%@1.0uM		
B-1206	81.0%@1.0uM	0.34uM		······································
B-1207	90.0%@1.0uM	58.0%@1.0uM	***	
B-1208	82.0%@1.0uM	51.0%@1.0uM		
B-1209	86.0%@1.QuM	55.0%@1.0uM		
B-1210	82.0%@1.0uM	57.0%@1.0uM		
B-1211	88.0%@1.0uM	59.0%@1.0uM		
B-1212	90.0%@1.0uM	57.0%@1.0uM		
B-1213	84.0%@1.0uM	0.62uM		
B-1214	76.0%@1.0uM	58.0%@1,0uM	·	
B-1215	86.0%@1.0uM	0.23uM		
B-1216	88.0%@1.0uM	0.18uM		
B-1217	87.0%@1.0uM	0.46uM		
B-1218	88.0%@1.0uM	76.0%@1.0uM		
B-1219	85.0%@1.0uM	37.0%@1.0uM		
3-1220	81.0%@1.0uM	53.0%@1.0uM		
3-1221	82.0%@1.0uM	44.0%@1.0uM		
3-1222	65.0%@1.0uM	9.0%@1.0uM		
3-1223	80.0%@1.0uM	61.0%@1.0uM		*****
3-1224	82.0%@1,0uM	74.0%@1.0uM		
3-1225	89.0%@1.0uM	73.0%@1.0uM		
3-1226	89.0%@1.0uM	0.18uM		
3-1227	83.0%@1.0uM	0.22uM		
3-1228	90.0%@1.0uM	0.72uM		
3-1229	87.0%@1.0uM	0.65uM		
3-1230	90.0%@1.0uM	0.25uM		
3-1231	94.0%@1.0uM	0.56uM		
3-1232	81.0%@1.0uM	54.0%@1.0uM		
3-1233	85.0%@1.0uM	0.36uM		
3-1234	89.0%@1.0uM	0.49uM		
3-1235	0.04uM	76.0%@1.0uM		

	P38 alpha kinase IC50,uM or %	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	inhib@conc. (uM)	or %	TNF inhib @ dose	inhib @dose
Example#	minowcone. (um)	inhib@conc. (uM)	@predose time	@predose time
B-1236	0.1uM	52 00/ 60 1 0 14		
B-1237	0.22uM	53.0%@1.0uM 39.0%@1.0uM		
B-1238	0.14uM			
B-1239	<0.1uM	16.0%@1.0uM		
B-1240	<0.1uM	38.0%@1.0uM		
B-1241	0.04uM	59.0%@1.0uM		
B-1242	0.08uM	81.0%@1.0uM		
B-1243	0.04uM	83.0%@1.0uM		
B-1244	0.26uM	47.0%@1.0uM		
B-1245	0.49uM	44.0%@1.0uM		
B-1246	0.49uM	42.0%@1.0uM		
B-1247	<0.1uM	40.0%@1.0uM		
B-1248	<0.1uM	58.0%@1.0uM		
B-1249	0.24uM	68.0%@1.0uM		
B-1250	0.14uM	60.0%@1.0uM		
B-1251	0.41uM	18.0%@1.0uM		
B-1252	0.17uM	38.0%@1.0uM		
B-1253		46.0%@1.0uM		
B-1254	0,15uM	57.0%@1.0uM		
B-1255	0.16uM	68.0%@1.0uM		
B-1256	12.9uM	75.0%@1.0uM		
B-1257	0.12uM	41.0%@1.0uM		
3-1258	1.48uM	40.0%@1.0uM		
3-1259	0.07uM	56.0%@1.0uM		
3-1259	<0.1uM	0.48uM		
3-1261	0.11uM	48.0%@1.0uM		
3-1262	0.74uM	44.0%@1.0uM		
3-1263	<0.1uM	63.0%@1.0uM		
3-1264	1.05uM	57.0%@1.0uM		
3-1265	0.32uM	47.0%@1.0uM		
-1266	0.43uM	51.0%@1.0uM		
3-1267	<0.1uM	58.0%@1.0uM		
1-1268	<0.1uM	73.0%@1.0uM		
-1269	<0.1uM	79.0%@1.0uM		
-1209	0.46uM	84.0%@1.0uM		
-1270	0.47uM	83.0%@1.0uM		
-1272	0.13uM	74.0%@1.0uM		
-1273	0.014uM	38.0%@1.0uM		
-1273	<0.1uM	36.0%@1.0uM		
-1274	<0.1uM	41.0%@1.0uM		
-1275 -1276	<0.1uM	50.0%@1.0uM		
-1276	0.062uM	11.0%@1.0uM		
-1277	<0.1uM	47.0%@1.0uM		
-1278	0.12uM	85.0%@1.0uM		
-1279	<0.1uM	79.0%@1.0uM		
	0.039uM	83.0%@1.0uM		
1281	<0.1uM	85.0%@1.0uM		
1282	<0.1uM	75.0%@1.0uM		
1283	<0.1uM	64.0%@1.0uM		
1284	<0.1uM	75.0%@1.0uM		

	P38 alpha kinase	U937 Cell IC50,uM	Mouse LPS Model %	Rat LPS Model %
	IC50,uM or %	or %	TNF inhib @ dose	inhib @dose
	inhib@conc. (uM)	inhib@conc. (uM)	@predose time	@predose time
Example#				
B-1285	0.057uM	80.0%@1.0uM		
B-1286	0.15uM	78.0%21.0uM		
B-1287	0.25uM	55.0%@1.0uM		
B-1288	0.15uM	74.0%@1.0uM		
B-1289	0.73uM	35.0%@1.0uM		
B-1290	0.26uM	75.0%@1.0uM		
B-1291	0.097uM	55.0%@1.0uM		
B-1292	0.01uM	74.0%@1.0uM		
B-1293	0.31uM	48.0%@1.0uM		
B-1294	0.013uM	54.0%@1.0uM		
B-1295	0.079uM	74.0%@1.0uM		······································
B-1296	0.038uM	48.0%@1.0uM		·
B-1297	0.02uM	>1.0uM		
B-1298	0.055uM	20.0%@1.0uM		
B-1299	0.091uM	>1.0uM		
B-1300	0.071uM	18.0%@1.0uM		
B-1301	0.12uM	15.0%@1.0uM		
B-1302	0.023uM	11.0%@1.0uM		
B-1303	0.08uM	>1.0uM		
B-1304	0,11uM	10.0%@1.0uM		
B-1305	0.64uM	9.0%@1.0uM	***********	
B-1306	0.11uM	>1.0uM		
B-1307	0.009uM	16.0%@1.0uM		
B-1308	<0.1uM	>1.0uM		
B-1309	0.045uM	>1.0uM		
B-1310	0.12uM	11.0%@1.0uM		
B-1311	0.05uM	57.0%@1.0uM		
B-1312	0.35uM	>1.0uM		
B-1313	0.035uM	37.0%@1.0uM		
B-1314	0.045uM	24.0%@1.0uM		
B-1315	0.055uM	12.0%@1.0uM		
3-1316	0.026uM	36.0%@1.0uM		
3-1317	0.019uM	9.0%@1.0uM		·
3-1318	<0.1uM	1.0%@1.0uM		
3-1319	0.24uM	>1.0uM		
3-1320	0.047uM	43.0%@1.0uM		
3-1321	0.47uM	66.0%@1.0uM		···
3-1322	0.12uM	87.0%@1.0uM		
3-1323	0.013uM	85.0%@1.0uM		
3-1324	0.16uM	83.0%@1.0uM		
3-1325	0.27uM	95.0%@1.0uM		
3-1326	0.092uM	84.0%@1.0uM		
3-1327	0.13uM	65.0%@1.0uM		
3-1328	0.032uM	86.0%@1.0uM		
3-1329	0.66uM	54.0%@1.0uM		
3-1330	0.053uM	85.0%@1.0uM		
3-1331	0.004uM	85.0%@1.0uM		
-1332	0.007uM	81.0%@1.0uM		
3-1333	0.45uM	76.0%@1.0uM		

	P38 alpha kinase IC50.uM or %	U937 Cell (C50,uM	Mouse LPS Model %	Rat LPS Model % inhib @dose
	inhib@conc. (uM)		@predose time	@predose time
Example#				
B-1334	0.13uM	73.0%@1.0uM		
B-1335	0.097uM	63.0%@1.0uM		
B-1336	0.072uM	83.0%@1.0uM		
B-1337	0.4uM	90.0%@1.0uM		
B-1338	0.18uM	73.0%@1.0uM		
B-1339	0.12uM	67.0%@1.0uM		
B-1340	0.043uM	63.0%@1.0uM		
B-1341	0.42uM	52.0%@1.0uM		
B-1342	0.25uM	59.0%@1.0uM		
B-1343	0.065uM	83.0%@1.0uM		
B-1344	0.014uM	86.0%@1.0uM		
B-1345	0.27uM	73.0%@1.0uM		
B-1346	0.043uM	86.0%@1.0uM		
B-1347	0.021uM	84.0%@1.0uM		
B-1348	0.009uM	69.0%@1.0uM		
B-1349	0.037uM	86.0%@1.0uM		
B-1350	0.019uM	78.0%@1.0uM		
B-1351	0.068uM	78.0%@1.0uM		
B-1352	0.013uM	76.0%@1.0uM		
B-1353	0.062uM	80.0%@1.0uM		*******
B-1354	0.013uM	83.0%@1.0uM		
B-1355	0.07uM	75.0%@1.0uM		
B-1356	0.059uM	91.0%@1.0uM		
B-1357	0.18uM	84.0%@1.0uM		
B-1358	0.16uM	76.0%@1.0uM		
B-1359	0.005	84.0%@1.0uM		
B-1360	0.11	0.15uM		54%@3mpk@-4h
B-1361	0.03	0.29uM		a compile in
B-1362	0.003	0.29uM		
B-1363	0.009	0.28uM	51.0%@30pmk @- 6H	53%@3mpk@-4h
B-1364	0.009	0.27uM	53.0%@30mpk@- 6.0H	17%@3mpk@-4h
B-1365	0.17	88.0%@1.0uM		
B-1366	0.04	0.27uM		
B-1367	<0.1	0.22uM		
B-1368	0.031	0.33uM	44.0%@30mpk @-	
B-1369	<0.1	0.29uM		
B-1370	<0.1	0.77uM		
B-1371	0.06	83.0%@1.0uM		
B-1372	<0.1	0.41uM	48.0%@30mpk @-	
B-1373	0.016	0.17uM		
B-1374	<0.1	0.28uM		
B-1375	0.01	0.25uM		
B-1376	0.009	0.26uM	3.0%@30mpk @-6H	
B-1377	0.12	5.0uM		
B-1378	0.02	1.04uM		
B-1379	<0.1	0.092uM		
B-1380	<0.1	0.26uM		

	P38 alpha kinase IC50,uM or %	U937 Cell 1C50,uM	Mouse LPS Model % TNF inhib @ dose	Rat LPS Model %
	inhib@conc. (uM)	inhlb@conc. (uM)	©predose time	@predose time
Example#				
B-1381	0.055	0.73uM		
B-1382	<0.1	0.44uM		
B-1383	0.0012	0.15uM		
B-1384	0.57	0.37uM		
B-1385	<0.1	0.11uM		
B-1386	<0.1	0.25uM		
B-1387	<0.1	0.1uM		
B-1388	0.57	1.38uM		
B-1389	0.06	0.57uM		
B-1390	<0.1	71.0%@1.0uM		
B-1391	0.016uM	82.0%@1.0uM		
B-1392	0.059uM	82.0%@1.0uM		
B-1393	3.17uM	80.0%@1.0uM		
B-1394	0.32uM	78.0%@1.0uM		
B-1395	1.48	61.0%@1.0uM		
B-1396	1.55	73.0%@1.0uM		
B-1397	0.92	85.0%@1.0uM		
B-1398	0.67	83.0%@1.0uM		
B-1399	0.14	74.0%@1.0uM		
B-1400	0.024	83.0%@1.0uM		
B-1401	0.033	75.0%@1.0uM		
B-1402	0.12	76.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
B-1403	4.54	71%@1.0uM		
B-1404	0.6	70%@1.0uM		
B-1405	0.28	70%@1.0uM		
B-1406	1.39	56.0%@1.0uM		
B-1407	0.4	71.0%@1.0uM		
B-1408	0,27	69.0%@1.0uM		
B-1409	<0.1	72.0%@1.0uM		
B-1410	<0.1	69%@1.0uM		
B-1411	<0.1	81.0%@1.0uM		
B-1412	0.097	80.0%@1.0uM		
B-1413	0.016	78.0%@1.0uM		
B-1414	0.025	83.0%@1.0uM		
B-1415	1.41	79.0%@1.0uM		
B-1416	0.14	81.0%@1.0uM		
3-1417	0.069	69.0%@1.0uM		
3-1418	1.01	82.0%@1.0uM		
3-1419	0.3	84.0%@1.0uM		
3-1420	<0.1	82.0%@1.0uM		
3-1421	0.014	75.0%@1.0uM		
3-1422	0.58	68.0%@1.0uM		
3-1423	1.58	84.0%@1.0uM		
3-1424	0.86	76.0%@1.0uM		
3-1425	0.09	83.0%@1.0uM		
3-1426	0.19	80.0%@1.0uM		
3-1427	<0.1	84.0%@1.0uM		
3-1428	<0.1	86.0%@1.0uM		
3-1429	<0.1	87.0%@1.0uM		

	P38 alpha kinase IC50,uM or % inhib@conc. (uM)	U937 Cell IC50,uM or %	Mouse LPS Model % TNF inhib @ dose	Rat LPS Model
Example#	minowconc. (um)	inhib@conc. (uM)	@predose time	@predose time
B-1430	0.75uM	35.0% @1.0uM		
B-1431	0.36uM	59.0% @1.0UM		
B-1432	0.11uM	58.0% @1.0uM		
B-1433	0.26uM	51.0% @1.0uM		
B-1434	0.19uM	21.0% @1.0uM		
B-1435	1.8uM	28.0% @1.0uM		
B-1436	1.0uM	45.0% @1.0uM		
B-1437	0.3uM	20.0% @1.0uM		
B-1438	2.01uM	23.0% @1.0uM		
B-1439	1.7uM	27.0% @1.0uM		
B-1440	0.87uM	17.0% @1.0uM		
B-1441	1.95uM	3.0% @1.0uM		
B-1442		66.0% @1.0uM		
B-1443	1.54uM	18.0% @ 1.0uM		
B-1444	0.014uM	83.0% @1.0uM		
B-1445	0.3uM	24.0% @1.0uM		
B-1446	0.43uM	27.0% @1.0uM		
3-1447	0.77uM	36.0% @1.0uM		
3-1448	0.5uM	34.0% @1.0uM		
3-1448	1.43uM	22.0% @1.0uM		
3-1450	1.61uM	50.0%@1.0uM		
3-1451	2.1uM	49.0%@1.0uM		· · · · · · · · · · · · · · · · · · ·
-1452	2.88uM	50% @1.0uM		
	2.41uM	47.0%@1.0uM		
3-1453	2.53uM	49.0% @1.0uM		
1-1454 1-1455	1.6uM	12.0% @1.0uM		
	1.21uM	8.0% @1.0uM		
-1456	1.29uM	>1.0uM		
-1457	0.43uM	43.0% @1.0uM		
-1458	0.95uM	65.0% @1.0uM		
-1459	0.67uM	46.0% @1.0uM		
-1460	0.96uM	29.0% @1.0uM		
-1461	0.4uM	39.0% @1.0uM		
-1462	0.22uM	50.0% @1.0uM		
1463	2.34uM	26.0% @1.0uM		
1464	1.18uM	27.0% @1.0uM		
1465	3.23uM	31.0% @1.0uM		
1466	1.69uM	>1.0uM		
1467	1.22uM	1.0% @1.0uM		
1468	1.61uM	10.0% @1.0uM		
1469	0.37uM	14.0% @1.0uM		
1470	0.6uM	28.0% @1.0uM		
1471	0.85uM	25.0% @1.0uM		
1472	0.93uM	12.0%@1.0uM		
1473	1.24uM	14.0% @ 1.0uM		
1474		31.0% @1.0uM		
1475		24.0% @1.0uM		
1476	0.047uM	12.0% @1.0uM		
1477		34.0% @1.0uM		
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Example#	P38 alpha kinase IC50,uM or % Inhib@conc. (uM)	or %	Mouse LPS Model % TNF Inhib @ dose @predose time	Rat LPS Model % inhlb @dose @predose time
B-1479				

Example#	iC50,uM or %	or %	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-2270	0.72uM	31%@10.0uM		
B-2271	0.93uM	38%@10.0uM		
B-2272	0.26uM	53.0%@10.0uM		
B-2273	1.92uM	39.0%@ 10.0uM		
B-2274	0.26uM	59.0%@10.0uM		
B-2275	2.16uM	53.0%@10.0uM		
B-2276	11.5uM	37.0%@10.0uM		
B-2277	14.9uM	44.0%@10.0uM		
B-2278	0.8uM	51.0%@10.0uM		
B-2279	0.32uM	36.0%@10.0uM		
B-2280	0.4uM	57.0%@10.0uM		
B-2281	0.81uM	60.0%@10.0uM		
B-2282	0.91uM	41.0%@10.0uM		
B-2283	0.04uM	53.0%@10.0uM		
B-2284	4.61uM	62.0%@10.0uM		
B-2285	2.29uM	49.0%@10.0uM		
B-2286	0.017uM	0.78uM	25%@30mpk@-1h	
B-2287	2.56uM	61.0%@10.0uM		
B-2288	6.51uM	46.0%@10.0uM		····
B-2289	3.0uM	30.0%@10.0uM		· · · · · · · · · · · · · · · · · · ·
B-2290	2.37uM	59.0%@10.0uM		
B-2291	0.019uM	41%@10.0uM		
B-2292	8.82uM	57.0%@10.0uM		
3-2293	2.11uM	56.0%@10.0uM		
3-2294	1.68uM	50.0%@10.0uM		
3-2295	1.79uM	56.0%@10.0uM		
3-2296	17.3uM	63.0%@10.0uM		
3-2297	3.59uM	57.0%@10.0uM		
3-2298	0.29uM	4.22uM		
3-2299	1.97uM	62.0%@10.0uM		
3-2300	0.07uM	43.0%@10.0uM		
3-2301	0.18uM	44.0%@10.0uM		
3-2302	1.0uM	58.0%@1.0uM		
3-2303	0.011uM	54.0%@10.0uM		***************************************
3-2304	1.41uM	50.0%@ 10.0uM		
-2305	0.54uM	60.0%@10.0uM		
-2306		39.0%@10.0uM		
-2307	2.29uM	69.0%@10.0uM		··········
-2308	0.66uM	56.0%@10.0uM		
-2309	0.29uM	47.0%@10.0uM		

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Example#	1C50,uM or %	or %	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-2310	0.12uM	1.2uM	50%@30mpk@-6h	
B-2311	7.18uM	60%@10.0uM	o o o o o o o o o o o o o o o o o o o	
B-2312	2.93uM	43.0%@10.0uM		
B-2313	42.3uM	58.0%@10.0uM		
B-2314	11.0uM	66.0%@10.0uM		
B-2315	0.49uM	36.0%@10.0uM		
B-2316	0.46uM	58.0%@10.0uM		
B-2317	1.0uM	60.0%@10.0uM		
B-2318	73.0%@10.0uM	25.0%@10.0uM		
B-2319	75.0%@10.0uM	40.0%@10.0uM	-	· · · · · · · · · · · · · · · · · · ·
B-2320	44.0%@10.0uM	35.0%@10.0uM		
B-2321	69.0%@10.0uM	27.0%@10.0uM		
B-2322	76.0%@10.0uM	38.0%@10.0uM		
B-2323	69.0%@10.0uM	46.0%@10.0uM		
B-2324	58.0%@10.0uM	36.0%@10.0uM		
B-2325	60.0%@10.0uM	51.0%@10.0uM		
B-2326	76.0%@10.0uM	33.0%@10.0uM		
B-2327	76.0%@10.0uM	23.0%@10.0uM		
B-2328	65.0%@10.0uM	28.0%@10.0uM		
B-2329	72.0%@10.0uM	53.0%@10.0uM		
B-2330	81.0%@10.0uM	37.0%@10.0uM		
B-2331	74.0%@10.0uM	44.0%@10.0uM		
B-2332	70.0%@10.0uM	47.0%@10.0uM		
B-2333	58.0%@10.0uM	36.0%@10.0uM		
B-2334	81.0%@10.0uM	45.0%@10.0uM		
B-2335	82.0%@10.0uM	50.0%@10.0uM		
B-2336	48.0%@10.0uM	35.0%@10.0uM		
B-2337	46.0%@10.0uM	59.0%@10.0uM		7
B-2338	73.0%@10.0uM	50.0%@10.0uM		
B-2339	84.0%@10.0uM	>10.0uM		
B-2340	35.0%@10.0uM	12.0%@10.0uM		
B-2341	75.0%@10.0uM	50.0%@10.0uM		
B-2342	83.0%@10.0uM	46.0%@10.0uM		
B-2343	43.0%@10.0uM	27.0%@10.0uM		
B-2344	71.0%@10.0uM	50.0%@10.0uM		
B-2345	64.0%@10.0uM	38.0%@10.0uM		
B-2346	45.0%@10.0uM	48.0%@10.0uM		
B-2347	49.0%@10.0uM	50.0%@10.0uM		
B-2348	76.0%@10.0uM	48.0%@10.0uM		•
B-2349	75.0%@10.0uM	27.0%@10.0uM		

Example#	IC50,uM or %	or %	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-2350	38.0%@10.0uM	56.0%@10.0uM		· · · · · · · · · · · · · · · · · · ·
B-2351	77.0%@10.0uM	1.0%@10.0uM		
B-2352	37.0%@10.0uM			
B-2353	38.0%@10.0uM	33.0%@10.0uM		
B-2354	65.0%@10.0uM			
B-2355		50.0%@10.0uM		
B-2356	77.0%@10.0uM	45.0%@10.0uM		 ,
B-2357	47.0%@10.0uM			
B-2358	17.0%@10.0uM			
B-2359	76.0%@10.0uM	35.0%@10.0uM		···
B-2360	45.0%@10.0uM	>10.0uM		
B-2361	19.0%@10.0uM			
B-2362	60%@100.0uM	39.0%@10.0uM		
B-2363	44.0%@10.0uM	1.0%@10.0uM		
B-2364	47.0%@10.0uM	4.0%@10.0uM		
B-2365	82.0%@10.0uM	43.0%@10.0uM		
B-2366	70.0%@10.0uM	59.0%@10.0uM		
B-2367	46.0%@10.0uM	40.0%@1.0uM		•
B-2368	65.0%@10.0uM	55.0%@10.0uM		
B-2369	32.0%@10.0uM	>10.0uM		
B-2370	73%@100.0uM	20.0%@10.0uM		
B-2371	54.0%@10.0uM	36.0%@10.0uM	· · · · · · · · · · · · · · · · · · ·	
B-2372	55.0%@100.0uM	>10.0uM		
B-2373	50.0%@100.0uM	6%@10.0uM		
B-2374	35.0%@10.0uM	20.0%@10.0uM		
B-2375	62.0%@100.0uM	>10.0uM		
B-2376	32.0%@10.0uM	17.0%@10.0uM		
B-2377	34.0%@10.0uM	17.0%@10.0uM		
B-2378	48.0%@10.0uM	61.0%@10.0uM		
B-2379	73.0%@100.0uM	45.0%@1.0uM		
B-2380	81%@100.0uM	53.0%@10.0uM		
B-2381	68%@100.0uM	2.0%@10.0uM		
B-2382	51.0%@10.0uM	24.0%@10.0uM		
B-2383	63.0%@10.0uM	35.0%@10.0uM		
B-2384	49%@100.0uM	10.0%@10.0uM		
B-2385	79.0%@10.0uM	19.0%@10.0uM		
B-2386	38.0%@10.0uM	19.0%@10.0uM		
B-2387	50.0%@100.0uM	>10.0uM		
B-2388	42.0%@10.0uM	24.0%@10.0uM		
B-2389	39.0%@10.0uM	29.0%@10.0uM		

Example#	P38 alpha kinase IC50,uM or %	U937 Ceil IC50,uM or %	Mouse LPS Model % TNF inhib @	Rat LPS Model % inhib @dose
Lyguibie			dose @predose time	@predose time
	,			
B-2390	34.0%@10.0uM	27.0%@1.0uM		
B-2391	40.0%@10.0uM	59.0%@10.0uM		
B-2392	63.0%@10.0uM	46.0%@10.0uM		
B-2393	43.0%@10.0uM	>10.0uM		
B-2394	37.0%@10.0uM	22.0%@10.0uM		
B-2395	32.0%@10.0uM	28.0%@10.0uM		
B-2396	75.0%@10.0uM	>10.0uM		
B-2397	83.0%@10.0uM	22.0%@10.0uM		•
B-2398	55%@100.0uM	10.0%@10.0uM		
B-2399	69.0%@10.0uM	18.0%@10.0uM		
B-2400	60.0%@10.0uM	40,0%@10.0uM		
B-2401	78.0%@10.0uM	44.0%@10.0uM		
B-2402	43.0%@10.0uM	52.0%@10.0uM		
B-2403	72%@100.0uM	52.0%@10.0uM		
B-2404	58%@100.0uM	52.0%@10.0uM		<u> </u>
B-2405	47%@100.0uM	>10.0uM		
B-2406	45.0%@10.0uM	24.0%@10.0uM		
B-2407	47%@100.0uM	27.0%@10.0uM		
B-2408	39.0%@10.0uM	10.0%@10.0uM		
B-2409	78.0%@10.0uM	26.0%@10.0uM		
B-2410	33.0%@10.0uM	32.0%@10.0uM		
B-2411	26%@100.0uM	13.0%@10.0uM		
B-2412	40.0%@10.0uM	31.0%@10.0uM		
B-2413	75.0%@10.0uM	37.0%@10.0uM		
B-2414	86.0%@10.0uM	38.0%@10.0uM		
B-2415	94.0%@10.0uM	50.0%@10.0uM		
B-2416	85.0%@10.0uM	43.0%@1.0uM		
B-2417	83.0%@10.0uM	18.0%@10.0uM		
B-2418	88.0%@10.0uM	34.0%@10.0uM		
B-2419	86.0%@10.0uM	66.0%@10.0uM		
B-2420	70.0%@10.0uM	34.0%@10.0uM		
B-2421	89.0%210.0uM	38.0%@10.0uM		
B-2422	90.0%@10.0uM	17.0%@10.0uM		
B-2423	85.0%@10.0uM	>10.0uM		
B-2424	86.0%@10.0uM	43.0%@10.0uM		
B-2425	79.0%@10.0uM	42.0%@10.0uM		
B-2426	88.0%@10.0uM	53.0%@10.0uM		
B-2427	87.0%@10.0uM	59.0%@10.0uM		
B-2428	82.0%@10.0uM	50.0%@10.0uM		
B-2429	92.0%@10.0uM	32.0%@10.0uM		

Example#	IC50,uM or %	or %	Mouse LPS Model % TNF inhib @ dose @predose time	Rat LPS Model % inhib @dose @predose time
B-2430	90.0%@10.0uM	61.0%@10.0uM		
B-2431	85.0%210.0uM	68.0%@10.0uM		
B-2432	86.0%210.0uM	40.0%@10.0uM		
B-2433	94.0%@10.0uM	84.0%@10.0uM		
B-2434	92.0%@10.0uM	63.0%@10.0uM		
B-2435	84.0%@10.0uM	4.0%@10.0uM		
B-2436	80.0%@10.0uM	54.0%@10.0uM		
B-2437	82.0%@10.0uM	41.0%@ 10.0uM		· · · · · · · · · · · · · · · · · · ·
B-2438	75.0%@10.0uM	40.0%@10.0uM		
B-2439	81.0%@10.0uM	44.0%@10.0uM		
B-2440	77.0%@10.0uM	78.0%@10.0uM		
B-2441	86.0%@10.0uM	46.0%@10.0uM		
B-2442	86.0%@10.0uM	>10.0uM		
B-2443	84.0%@10.0uM	44.0%@10.0uM		
B-2444	89.0%@10.0uM	7.0%@10.0uM		
B-2445	94.0%@10.0uM	15.0%@10.0uM		
B-2446	90.0%@10.0uM	28.0%@10.0uM		
B-2447	94.0%@10.0uM	>10.0uM		
B-2448	75.0%@10.0uM	30.0%@10.0uM		
B-2449	86.0%@10.0uM	42.0%@10.0uM		
B-2450	87.0%@10.0uM	46.0%@1.0uM		
B-2451	87.0%@10.0uM	45.0%@10.0uM		
B-2452	89.0%@10.0uM	33.0%@10.0uM		
B-2453	91.0%@10.0uM	>10.0uM		
B-2454	88.0%@10.0uM	40.0%@10.0uM		
B-2455	87.0%@10.0uM	54.0%@10.0uM		
B-2456	86.0%@10.0uM	53.0%@10.0uM		
B-2457	90.0%@10.0uM	18.0%@10.0uM		
B-2458	83.0%@10.0uM	36.0%@10.0uM		*
B-2459	82.0%@10.0uM	81.0%@10.0uM		
B-2460	80.0%@10.0uM	79.0%@10.0uM		
B-2461		59.0%@10.0uM		
			-	

Biological data from a number of compounds of Examples C-74 through C-139 are shown in the following tables.

In vitro P38-alpha kinase inhibitory data are shown in the column identified as:

"P38 alpha kinase IC50, µM"

In vitro human whole blood assay data for measuring the ability of the compounds to inhibit TNF production in human whole blood stimulated with LPS are shown in the column identified as:

"Human Whole Blood IC50, µM or %Inhib@conc. (µM)"

In vivo assessment of the ability of the compounds to inhibit LPS-stimulated TNF release in the rat is shown in the column identified as:

"Rat LPS Model % Inhibition@dose@predose time"
wherin the dose is milligram per kilogram (mpk)
administered by oral gavage and the predose time
indicates the number of hours before LPS challenge when
the compound is administered.

Example#	IC50, μM	Human Whole Blood IC50, µM or %Inhib@conc. (µM)	Rat LPS Model % Inhibition@ dose@predose time
C-74	0.037	0.56	54%@5mpk@-4h
C-75	0.045	0.4	71%@5mpk@-4h
C-76	0.07	3.24	66%@5mpk@-4h
C-77	0.071	8.2	92%@5mpk@-4h
C-78	0.068	10.5	87%@5mpk@-4h
C-79	0.045	0.52	83%@5mpk@-4h

Example#	P38 alpha kinase	Human Whole Blood	Rat LPS Model
1	IC50, μM	IC50, µM or	% Inhibition@
		%Inhib@conc. (µM)	dose@predose
			time
C-80	0.008	51%@ 5 μM	
C-81	0.037	40%@ 5 μM	
C-82	0.15	7.31	
C-83	0.24	1.23	25%@5mpk@-4h
C-84	0.048	0.88	22%@5mpk@-4h
C-85	0.57	>25	
C-86	0.007	0.19	66%@5mpk@-4h
C-87	0.027	0.34	
C-88	0.012	0.3	59%@5mpk@-4h
C-89	0.039	0.12	27%@5mpk@-4h
C-90	0.037	0.48	
C-91	0.054	2.31	63%@5mpk@-4h
C-92	0.024	0.28	66%@5mpk@-4h
C-93	0.009	0.38	50%@5mpk@-4h
C-94	0.02	0.27	73%@5mpk@-4h
C-95	0.13	3.91	32%@5mpk@-4h
C-96	0.077	2.1	38%@5mpk@-4h
C-97	0.025	3.83	21%@5mpk@-4h
C-98	0.016	0.64	78%@5mpk@-4h
C-99	0.062	0.38	36%@5mpk@-4h
C-100	0.027	0.27	44%@5mpk@-4h
C-101	0.083	3.71	52%@5mpk@-4h
C-102	0.29	7.56	72%@5mpk@-4h
C-105	0.033	0.13	46%@5mpk@-4h
C-106	0.026	0.44	23%@5mpk@-4h
C-107	0.014	0.38	11%@5mpk@-4h
C-108	0.02	0.73	0%@5mpk@-4h
C-111	0.21	6.05	39%@5mpk@-4h
C-112	0.54	6.36	89%@5mpk@-4h
C-113	0.082	2.72	77%@5mpk@-4h
C-114	0.11	1.73	39%@5mpk@-4h
C-115	0.042	10.2	39%@5mpk@-4h
C-116	0.429	0.50	53%@5mpk@-4h
C-117	3.42	7.26	71%@5mpk@-4h
C-118	0.298	>25	39%@5mpk@-4h
C-120	0.7	18.6	26%@5mpk@-4h
C-121	0.11	15.3	39%@5mpk@-4h
C-122	0.025		55%@5mpk@-4h
C-123	0.67	>25.0	

Example#	P38 alpha kinase IC50, μΜ	Human Whole Blood IC50, µM or %Inhib@conc. (µM)	Rat LPS Model % Inhibition@ dose@predose time
C-124	0.17	4.56	51%@20mpk@-4h
C-125	7.22	>25.0	
C-126	0.71	>25.0	6%@20mpk@-4h
C-127	0.038	0.27	53%@5mpk@-4h
C-128	0.09	2.22	63%@5mpk@-4h
C-132	0.086	44%@ 5 µM	
C-133	0.16	4.54	55%@5mpk@-4h
C-135	6.0		
C-136	0.032		
C-137	0.051		58%@5mpk@-4h
C-138	0.28	0.68	26%@5mpk@-4h
C-139	0.2	3.66	46%@5mpk@-4h

C-3015/2

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Additional compounds of interest can be prepared as set forth above and as described below in Scheme D-1, wherein the R_1 and R_2 substituents are as defined previously.

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The synthesis begins with the treatment of 4-methylpyrimidine 2 with a base such as LiHMDS, LDA or tBuOK in an organic solvent such as THF or ether which is cooled in an ice bath (0-10 °C). To the resulting 4-methylanion is added a solution of a suitably protected (Boc is shown) ethyl ester of isonipecotic acid 1 in THF or ether. The reaction is allowed to warm to room

1055

temperature and stirred for a period of 4 hours to 20 hours at which time the desired ketone 3 is isolated after aqueous work up. Condensation of the ketone 3 with tosylhydrazide in toluene or benzene as a solvent refluxing temperatures for a period of 1 hour to 5 hours affords the hydrazone 4. The hydrazone 4 is reacted with a suitably substituted benzoyl chloride 5, in the presence of a base such as LiHMDS or LDA or tBuOK or triethylamine at temperatures ranging from 0 °C to 70 °C. The reaction 10 is stirred for a period of 3-6 hours. Acidic hydrolysis of the protecting groups with an aqueous acid such as HCl or H2SO, and subsequent neutralization with an aqueous base such as NaOH or KOH affords the desired pyrazole 6. Treatment of the pyrazole 6 with an acid chloride 7 in the presence of base or with an acid 8 under standard peptide coupling conditions (EDC or DCC or PyBrOP with an additive such as HOBt or HATU and base such as N-methylmorpholine or diisopropylethylamine or triethylamine) affords the desired pyrazole amide 9. In most instance the desired products can be obtained pure by direct trituration with solvents such as methanol, ethyl acetate, acetonitrile or ether and/or recrystallization from suitable solvents.

The following examples contain detailed descriptions of methods of preparation of these additional compounds that form part of the invention. These detailed descriptions are presented for illustrative purposes only and are not intended as a restriction on the scope of the invention. All compounds showed NMR spectra consistent with their assigned structures.

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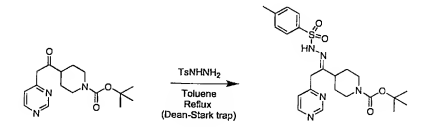
N-(2-Hydroxyacety1)-5-(4-piperidy1)-4-(4-pyrimidy1)-3-(4-chloropheny1)pyrazole

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Step 1: A 5 L 4-necked round bottom flask fitted with an overhead mechanical stirrer, N₂ inlet and a thermocouple was charged with 600 g (2.75 mol) of di-tert-butyl-dicarbonate and 1.5 L of CH₂Cl₂. The solution was cooled to 0 °C and 428 g (2.73 mol) of ethyl isonipecotate was added dropwise via an addition funnel. The addition took 45 minutes and the temperature rose from 0 °C to 17.4 °C. The reaction mixture was stirred for an additional 2 hours at ambient temperature. The solvent was removed in vacuo to afford 725 g of a yellow oil (residual solvent remained).

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Step 2: A 3 L 3-necked round bottom flask fitted with an overhead mechanical stirrer, a N_2 inlet, an addition 5 funnel and a thermocouple was charged with 1850 mL (1.85 mol) of a 1.0 M solution of LiHMDS in THF. The flask was cooled to 5 °C and 68 mL (0.74 mol) of 4-methylpyrimidine was added (neat) to the stirred solution. To this solution was added 198 g (0.77 mol) of Ethyl-N-t-10 butylcarbonyl isonipecotate dissolved in 160 mL of THF. The ice bath was removed and the reaction was allowed to stir for 18 hours. The reaction was quenched with 500 mL of saturated NH₄Cl and was extracted with 500 mL of ethyl acetate. The organic phase was washed with 500 $\ensuremath{\text{mL}}$ of 15 brine, dried over anhydrous Na₂SO₄, filtered concentrated in vacuo to afford 235 g of a brown oil.



20 Step 3: A 2 L 3-necked round bottom flask fitted with an overhead mechanical stirrer, a Dean-Stark trap and

a thermocouple was charged with 1.5 L of toluene, 226 q (0.742 mol) of N-t-butylcarbonyl-1-(4-piperidyl)-2-(4pyrimidyl)-1-ethanone and 138.4 g (0.743 mol) of tosyl hydrazide. The mixture was warmed to reflux. solution was allowed to reflux for 2 hours and was cooled 5 to ambient temperature. The reaction was allowed to stand overnight. A fine precipitate formed and was removed by filtration. The filtrate was concentrated in vacuo to afford a brown solid. The solid was suspended in 500 mL of ethyl acetate and the resulting mixture was placed in a sonication bath for 5 hours. The mixture was cooled in an ice bath and was filtered to afford 310 g of a wet solid. The solid was dried in a vacuum oven (40 °C, 5 mm) overnight to afford 248 g of the desired hydrazone (71%). ¹H NMR (CDCl₃) δ 9.03 (d, J = 1.2 Hz, 1H), 8.72 (d, J = 5.2 Hz, 2H), 7.89 (d, J = 8.3 Hz, 2H), 7.32 (d, J = 8.1 Hz, 2H), 7.26 (dd, J = 5.2, 1.0 Hz, 1H), 4.03 (d, J = 12.1 Hz, 2H), 3.76 (s, 2H), 2.71 (t, J = 12.1 Hz, 2H), 2.43 (s, 3H), 2.34 (m, 1H), 1.66 (d, J = 13.5 Hz, 2H), 1.47 (s, 9H), 1.38 (m, 2H); MS (M + H): 474 (base peak).

Step 4:

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Method A. A 2 L 3-necked round bottom flask fitted with an overhead mechanical stirrer, a N_2 inlet, an addition funnel and a thermocouple was charged with 400 mL (400 mmol) of a 1.0 M solution of LiHMDS in THF. solution was cooled to -21.9 $^{\circ}\text{C}$ and a solution of 62 g 5 of N-t-butylcarbonyl-1-(4-piperidyl)-2-(4pyrimidyl)-1-ethanone p-toluenesulfonyl hydrazone in 400 mL of THF was added slowly. The temperature never exceeded -11 °C throughout the addition. The solution was re-cooled to -19.6 $^{\circ}\text{C}$ and 23.0 g (131 mmol in 250 mL of 10 THF) of p-chlorobenzoylchloride was added slowly. temperature never exceeded -13 °C throughout the addition. The cooling bath was removed and the reaction was allowed to warm to ambient temperature. After 3 hours the reaction was quenched with 600 mL of 3 N HCl. The reaction was warmed to reflux and was held at reflux for 2 The reaction was allowed to cool to ambient temperature overnight. The reaction mixture was washed with 1.4 L of Et_2O and the aqueous phase was neutralized with 1 L of 2.5 N NaOH. The aqueous phase was extracted 20 with ethyl acetate (2 x 1000 mL). The combined organic phases were washed with brine (1 \times 500 mL), dried over anhydrous Na_2SO_4 , filtered and concentrated in vacuo to afford 21 g of a yellow solid. The solid was suspended in 500 mL of 2:1 $\mathrm{Et}_2\mathrm{O}/\mathrm{hexane}$. After sonication the solid was 25 isolated by filtration to leave a wet solid. The solid was dried in a vacuum oven to afford 13.8 g of 5-(4piperidyl) -4-(4-pyrimidyl) -3-(4-chlorophenyl) ¹H NMR (DMSO- d_6) 9.18 (s, 1H), 8.65 (d, J = 5.2, 1H), 7.44 (d, J = 8.5, 2H), 7.37 (d, J = 7.7 Hz, 2H), 7.15 (d, 30

1060

J = 5.2 Hz, 1H), 3.16 (m, 1H), 3.00 (d, J = 11.9 Hz, 2H), 2.52 (m, 2H), 1.69 (m, 4H); MS (M + H): 340 (base peak).

Method B: To a solution of 200 g (423 mmol) of N-tbutylcarbonyl-1-(4-piperidyl)-2-(4-pyrimidyl)-1-ethanone p-toluenesulfonyl hydrazone in 800 πL THF was added 70 πL 10 (500 mmol) of triethylamine in a 3 L three necked flask. The solution was cooled in an ice/salt/water bath to 0-5 To this cold solution was added a solution of 4chlorobenzoyl chloride (74 g, 423 mmol) in 100 mL THF 15 dropwise, maintaining the temperature below 10 °C. After the addition was complete the ice-bath was removed and replaced with a heating mantle. 4-N, Ndimethylaminopyridine (5 g, 40 mmol) was added and the reaction mixture was heated to 50 °C for 15-30 minutes. The reaction mixture was filtered and the residue washed

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1061

with THF (100 mL). The combined filtrates were evaporated under reduced pressure to a semisolid.

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The semisolid residue was dissolved in 450 mL THF and 180 mL of 12 N HCl was added to this solution rapidly. The reaction mixture was heated to 65 °C for 1.5-2 hours and transferred to a separatory funnel. The organic layer was discarded and the aqueous phase was washed twice with 200 mL of THF. The aqueous phase was transferred back to a 2 L flask and cooled to 0-10 °C in an ice bath. of the solution was adjusted to between ~ 9-10 by dropwise addition of 15 N ammonium hydroxide (~ 180 mL). mixture was transferred back to a separatory funnel and extracted with warm n-butanol (3 X 150 mL). The combined n-butanol phases were evaporated under reduced pressure to dryness. The residue was then stirred with methanol (200 mL), filtered and dried to obtain 129 g (90%) of the desired 5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl) pyrazole as a off-white solid. This material was identical in all respects to the material prepared by Method A.

25 Step 5: A 1 L round bottom flask was charged with 34.2 g (102 mmol) of 5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl) pyrazole, 500 mL of CH₂Cl₂ and 26.6 mL (153 mmol) of Hunig's base. To this suspension was added 16.5

1062

g (122 mmol) of 1-hydroxybenzotriazole and 8.1 g (106 mmol) of glycolic acid. The addition of glycolic acid was followed by the addition of 23.7 g (122 mmol) of 1-(3dimethylaminopropyl)-3-ethylcarbodiimide hydrochloride. The reaction was allowed to stir at ambient temperature The reaction was concentrated in vacuo to overnight. leave an oily residue. The residue was dissolved in 400 mL of methanol and 50 mL of 2.5 N NaOH. The reaction mixture was stirred at ambient temperature for 1 hour. 10 The mixture was acidified to pH 5 with 2 N HCl and was extracted with CH,Cl, (6 \times 200 mL). The combined organic phases were filtered through phase paper and the filtrate was concentrated in vacuo to leave a vellow residue. The residue was treated with 75 mL of acetonitrile. Α 15 precipitate formed. The solid was filtered and washed with additional acetonitrile and Et₂O to afford 31.4 g of N-(2-hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4chlorophenyl) pyrazole. ¹H NMR (DMSO-d₆) 9.20 (s. 1H). 8.67 (d, J = 4.8, 1H), 7.40 (m, 4H), 7.17 (d, J = 4.0, 20 1H), 4.53 (m, 2H), 4.13 (s, 2H), 3.77 (m, 1H), 3.05 (t, J= 12.7 Hz, 1H), 2.69 (m, 1H), 1.90 (m, 2H), 1.73 (m, 2H); MS (M +H): 398 (base peak).

Example D-2

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N-(2-Hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole hydrochloride

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A 25 mL round bottom flask was charged with 65 mg 5 (0.164 mmol) of N-(2-hydroxyacetyl)-5-(4-piperidyl)-4-(4pyrimidyl)-3-(4-chlorophenyl) pyrazole and 2.5 mL of dioxane. To this suspension was added 0.082 mL of 4 N HCl in dioxane. The mixture was stirred for 2 hours. The mixture was diluted with 5 mL of Et,0 and filtered. solid was dried over solid CaSO, under vacuum for 12 h to 10 afford 68 mg of N-(2-hydroxyacetyl)-5-(4-piperidyl)-4-(4pyrimidyl) -3-(4-chlorophenyl) pyrazole hydrochloride. 1H NMR (DMSO-d₄) 9.18(s, 1H), 8.63(d, J=5.37 Hz, 1H), 7.40 (d, J=8.59 Hz, 2H), 7.33 (d, J=8.59 Hz, 2H), 7.15 (m, 15 1H), 4.40(m, 1H), 4.06(m, 2H), 3.72(m, 1H), 3.33(m, 1H), 2.97(m, 1H), 2.62(m, 1H), 1.83(m, 2H), 1.64(m, 2H); MS (M+H): 398

Example D-3

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N-(2-Methoxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4chlorophenyl)pyrazole (fumarate salt)

1064

To a suspension of 250 mg (0.74 mmol) of 5-(4piperidyl) -4-(4-pyrimidyl) -3-(4-chlorophenyl) 5 (Example C-1, Step 3) and 180 mg (1.48 mmol) of N.Ndimethylamino pyridine in 20 mL of CH2Cl, was added 88 mg (0.81 mmol) of 2-methoxyacetyl chloride. The reaction was stirred for 5 hours. The reaction was quenched with 20 mL of saturated NH,Cl. The mixture was extracted with n-10 butyl alcohol and the organic layer was washed with brine. The solvent was removed to afford 72 mg of an oil. This oil was dissolved in 1 mL of warm MeOH. This solution was combined with a warm solution of 1 equivalent of fumaric acid in warm MeOH. The solution was cooled to ambient 15 temperature and the reaction was allowed to stir for 1 The solvent was removed in vacuo and the residue was triturated with Et₂O. The resulting solid was isolated by filtration to yield 56 mg of an off-white powder. 1 H NMR (DMSO- d_{6}) 13.23 (bs, 1H), 9.19 (d, J =1.2 Hz, 1H), 8.65 (d, J = 5.1 Hz, 1H), 7.41 (m, 4H), 7.16 20 (dd, J = 5.4, 1.2 Hz, 1H), 4.45 (bd, J = 11.1 Hz, 1H), $4.11 (q_{AB}, J = 39.0, 13.8 Hz, 2H), 3.86 (bd, J = 12.9 Hz,$ 1H), 3.32 (m, 4H), 3.04 (bt, J = 12.3 Hz, 1H), 2.63 (bt, J= 12.0 Hz, 1H), 1.77 (m, 4H); MS (M + H): 411 (base 25 peak).

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Example D-4

N-(2-Hydroxy-2-methylpropionyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole hydrochloride

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Step 1: To a suspension of 2.05 g (6.1 mmol) of 5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl) pyrazole 10 (Example C-1, Step 3) and 3.7 g (30.5 mmol) of N.Ndimethylamino pyridine in 30 mL of CH2Cl2 was added 1.06 mL (7.3 mmol) of 2-acetoxy-2-methylpropionyl chloride. The reaction was allowed to stir overnight at ambient temperature. The reaction was quenched with saturated 15 NH4Cl and water. The resulting aqueous phase was extracted with CH,Cl2. The combined organic layers were concentrated in vacuo to leave an oily solid. The residue was treated with CH₁CN and allowed to stand for 15 minutes. The resulting suspension was diluted with Et20 20 and was filtered to afford 2.2 g of a solid. Analysis by LC/MS indicated that the solid was a mixture of the hydroxy derivative and the acetoxy derivative. This solid was carried on to the next step without further purification.

Step 2: A solution of 1 g of the solid from step 1 in 10 mL of MeOH was treated with 500 mg of solid K_2CO_3 . The mixture was allowed to stir overnight at ambient

1066

temperature. The suspension was treated with water and the resulting solution was extracted with ethyl acetate. The organic phase was filtered through phase separation paper (to remove the residual water) and was concentrated The solid was dried in vacuo to leave an oily solid. under vacuum and was treated with CH,CN. The suspension was filtered to afford 825 mg of an off-white solid. solid was suspended in 5 mL of dioxane and 0.5 mL of 4 N HCl in dioxane was added. The suspension was stirred for 10 1 hour and the suspension was filtered to leave a solid. solid was washed with Et,O and the resulting suspension was filtered to give 900 mg of the title compound. H NMR (DMSO-d₆) 9.23 (s, 1H), 8.69 (s, 1H), 7.45 (m, 4H), 7.19 (s, 1H), 4.8 (br m, 4H), 3.85 (m, 2H). 15 3.38 (m, 1H), 1.89 (m, 2H), 1.72 (m, 2H), 1.37 (s, 6H); MS (M + H): 426 (base peak).

Example D-5

20 (S)-N-(2-Hydroxypropionyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole hydrochloride

25 By following the method of Example C-1 and substituting (S)-lactic acid for glycolic acid the title compound was prepared. ¹H NMR (DMSO-d₆) 13.15(s, br, 1H), 9.12(d, J=1.07 Hz, 1H), 8.59(d, J=5.37Hz, 1H),

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7.39(d, J=7.79Hz, 2H), 7.31(d, J=8.33, 2H), 7.10(dd, J=1.34, 5.1Hz, 1H), 4.76(m, 1H), 4.41(m, 2H), 3.99(m, 1H), 2.97(m, 1H), 2.45(m, 1H), 1.83(m, 2H), 1.64(m, 2H). 1.15(m, 3H); MS (M+H): 412 (base peak).

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Example D-6

(R)-N-(2-Hydroxypropionyl)-5-(4-piperidyl)-4-(4pyrimidyl)-3-(4-chlorophenyl)pyrazole hydrochloride

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Ву following the method of Example C-1 substituting (R)-lactic acid for glycolic acid the title compound was prepared. ¹H NMR (CDCl₁) 15 9.24(s, 1H), 8.52(d, J = 5.0 Hz, 1H), 7.32-7.36(m, 4H), 6.98(d, J = 5.3)Hz, 1H), 4.72 (d, J = 10.5 Hz, 1H), 4.55 (br, 1H), 3.88 (d, J= 13.1 Hz, 1H), 3.66(br, 1H), 3.19(br, 1H), 2.82(t, J =12.4 Hz, 1H), 2.10(br, 2H), 1.37(d, J = 6.2 Hz, 3H), 1.81-1.90(m, 2H); MS (M + H): 412 (base peak).

Example D-7

(R)-N-(2-Hydroxy-2-phenylacetyl)-5-(4-piperidyl)-4-(4pyrimidyl) -3-(4-chlorophenyl) pyrazole

1068

By following the method of Example C-1 and substituting (R)-phenylacetic acid for glycolic acid the title compound was prepared. ^{1}H NMR (DMSO- d_{6}) 9.15 (d, J=0.9 Hz, 1H), 8.63 (d, J=5.4 Hz, 1H), 7.40 (m, 9H), 7.13 (t, J=6.6 Hz, 1H), 5.43 (d, J=19.5 Hz, 1H), 4.51 (s, 1H), 4.04 (m, 1H), 3.33 (m, 4H), 2.8 (m, 2H), 1.68 (m, 3H); MS (M + H): 474 (base peak).

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Example D-8

N-(2-Hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4-fluorophenyl)pyrazole

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By following the method of Example C-1 and substituting 4-fluorobenzoyl chloride for 4-chlorobenzoyl chloride the title compound was prepared. ¹H NMR (DMF-d₇) 13.48(s, 1H), 9.40(s, 1H), 8.86(d, J = 5.1 Hz, 1H), 7.71(br, 2H), 7.42(bd, J = 5.2 Hz, 3H), 4.78(br, 1H), 4.43(s, 2H), 4.04(br, 1H), 3.79(br, 1H), 3.70(s, 1H),

1069

3.34(t, J = 12.2 Hz, 1H), 3.0(br, 1H), 2.21(d, J = 10.9 Hz, 2H), 2.08(br, 1H); MS (M + H): 382 (base peak).

Example D-9

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N-(2-Hydroxyacety1)-5-(4-piperidy1)-4-(4-pyrimidy1)-3-(4-trifluoromethylpheny1)pyrazole

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By following the method of Example C-1 and substituting 4-trifluoromethylbenzoyl chloride for 4-chlorobenzoyl chloride the title compound was prepared.

¹H NMR (DMF-d₇) 13.47(s, 1H), 9.24(s, 1H), 8.73(d, J = 4.0 Hz, 1H), 7.77(bd, J = 13.3 Hz, 4H), 7.34(d, J = 4.3 Hz, 1H), 4.61(br, 1H), 4.26(s, 2H), 3.87(br, 1H), 3.52(s, 2H), 3.17(t, J = 12.0 Hz, 1H), 2.8 (br, 1H), 2.02(br, 2H), 1.91(br, 1H); MS (M + H): 432 (base peak).

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Example D-10

N-(2-Hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(4trifluoromethoxyphenyl)pyrazole

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By following the methodof Example C-1 substituting 4-trifluoromethoxybenzoyl chloride for 4-5 chlorobenzoyl chloride the title compound was prepared. ¹H NMR (DMF-d₂) 13.55(s, 1H), 9.40(s, 1H), 8.88(d, J =4.6 Hz, 1H), 7.81 (d, J = 7.7 Hz, 2H), 7.64 (br, 2H), 7.47(d, J = 4.4 Hz, 1H), 4.75(br, 1H), 4.42(s, 2H),4.04(d, J = 12.5 Hz, 1H), 3.69(br, 2H), 3.34(t, J = 12.0)Hz, 1H), 3.0(br, 1H), 2.20(d, J = 11.7 Hz, 2H), 2.05(br. 1H); MS (M + H): 448 (base peak).

Example D-11

15 N-(2-Hydroxyacety1)-5-(4-piperidy1)-4-(4-pyrimidy1)-3-(3chlorophenyl)pyrazole

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20 By following the method of Example C-1 substituting 3-chlorobenzoyl chloride for 4-chlorobenzoyl chloride the title compound was prepared. H NMR (DMF-d₂) 13.41(s, 1H), 9.24(s, 1H), 8.73(d, J = 4.9 Hz, 1H), 7.56(s, 1H), 7.49(br, 2H), 7.41(br, 1H), 7.32(d, J = 4.2)

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Hz, 1H), 4.60(d, J = 11.7 Hz, 1H), 4.25(s, 2H), 3.87(d, J = 12.7 Hz, 1H), 3.52(bs, 2H), 3.17(t, J = 12.1 Hz, 1H), 2.84(d, J = 12.5 Hz, 1H), 2.03(d, J = 11.9 Hz, 2H), 1.87(br, 1H); MS (M + H): 398 (base peak).

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Example D-12

N-(2-Hydroxyacetyl)-5-(4-piperidyl)-4-(4-pyrimidyl)-3-(3-fluorophenyl)pyrazole

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By following the method of Example C-1 and substituting 3-fluorobenzoyl chloride for 4-chlorobenzoyl chloride the title compound was prepared. ¹H NMR (DMF-d₇) 13.38(s, 1H), 9.24(s, 1H), 8.72(d, J = 5.2 Hz, 1H), 7.49(dd, J = 8.0 and 6.2 Hz, 1H), 7.24-7.32(m, 4H), 4.60(d, J = 13.1 Hz, 1H), 4.25(s, 2H), 3.87(d, J = 13.3 Hz, 1H), 3.55-3.60(m, 1H), 3.52(s, 1H), 3.17(t, J = 12.2 Hz, 1H), 2.82(d, J = 12.9 Hz, 1H), 2.03(d, J = 10.9 Hz, 2H), 1.83-1.96(m, 1H); MS (M + H): 382 (base peak).

Example D-13

N-(2-Hydroxyacety1)-5-(4-piperidy1)-4-(4-pyrimidy1)-3-(3-trifluoromethylpheny1)pyrazole

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By following the method of Example C-1 substituting 3-trifluoromethylbenzoyl chloride for 4-5 chlorobenzoyl chloride the title compound was prepared. ¹H NMR (DMF-d₂) 13.76(s, 1H), 9.41(s, 1H), 8.91(d, J =5.3 Hz, 1H), 8.02(s, 1H), 7.95(t, J = 6.5 Hz, 2H), 7.85(t, J = 7.5 Hz, 1H), 7.53(d, J = 4.6 Hz, 1H), 4.78(d, J = 11.9)Hz, 1H), 4.45 (d, J = 16.3 Hz, 2H), 4.06 (d, J = 12.5 Hz, 10 1H), 3.69(bs, 2H), 3.34(t, J = 11.3 Hz, 1H), 3.01(d, J =13.1 Hz, 1H), 2.20(d, J = 11.1 Hz, 2H), 2.12(br, 1H); MS (M + H): 432 (base peak).

The following examples can be prepared in a manner similar to that described above for the synthesis of Examples C1-C13.

Example D-14

20 5-[4-N-(2-hydroxy-2-(2-chlorophenyl)acetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

1073

Example D-15

5-[4-N-(2-hydroxy-2-(3-chlorophenyl)acetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-16

5 = [4-N-(1-hydroxy-1-cyclohexylacetyl)piperidyl] -4-(4pyrimidyl) -3-(4-chlorophenyl)pyrazole

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Example D-17

5-[4-N-(2-hydroxy-1-cyclohexylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

1074

Example D-18

5-[4-N-(3-hydroxy-1-cyclohexylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

Example D-19

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5-[4-N-(4-hydroxy-1-cyclohexylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-20

5-[4-N-(1-hydroxy-1-cyclopentylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

1075

Example D-21

5-[4-N-(2-hydroxy-1-cyclopentylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

Example D-22

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5-[4-N-(3-hydroxy-1-cyclopentylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-23

5-[4-N-(3-hydroxypropionyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

1076

Example D-24

5-[4-N-(2-hydroxy-3,3,3-trifluoropropionyl)piperidyl]-4(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

Example D-25

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5-[4-N-(2-hydroxy-3-methylbutyryl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-26

5-[4-N-(2-hydroxyisocaproy1)piperidy1]-4-(4-pyrimidy1)-3-(4-chloropheny1)pyrazole

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Example D-27

5-[4-N-(2-hydroxy-2-cyclohexylacetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

Example D-28

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5-[4-N-(2-hydroxy-2-(4-methoxyphenyl)acetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-29

5-[4-N-(2-hydroxy-2-(3-methoxyphenyl)acetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-30

5 5-[4-N-(2-hydroxy-2-(4-trifluoromethylphenyl)acetyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-31

5-[4-N-(2-hydroxy-3-phenylpropionyl)piperidyl]-4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

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Example D-32

5-[4-N-(2-hydroxy-3-(4-hydroxyphenyl)propionyl)piperidyl}4-(4-pyrimidyl)-3-(4-chlorophenyl)pyrazole

N-NH N-ON

5-[4-N-(2-hydroxy-3-imidazolpropionyl)piperidyl]-4-(4pyrimidyl)-3-(4-chlorophenyl)pyrazole

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The synthesis of 2-substituted pyrimidinyl pyrazoles is shown in Scheme 2. Reaction of 2-methylmercapto-4methyl pyrimidine 10 with N-Boc methvl ester of isonipecotic acid (1) under basic (base selected from LiHMDS or LDA or tBuOK) conditions in an anhydrous solvent such as tetrahydrofuran or ether affords the desired Condensation of the ketone 11 with tosyl ketone 11. hydrazine under refluxing conditions in either toluene or

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benzene affords the hydrazone 12. The hydrazone 12 is deprotonated under basic (base selected from LiHMDS or LDA or tBuOK) conditions in an anhydrous solvent such as tetrahydrofuran or ether and the anion is reacted in situ with a suitably substituted benzoyl chloride 5 to afford, after mild aqueous work up, the desired and fully protected pyrazole 13. Oxidation of the 2-mercaptomethyl group present in 13 with oxidants selected from but not limited to Oxone, H,O, or mCPBA in solvents such as dichloromethane, acetonitrile or tetrahyrofuran affords the 2-methane sulfonyl pyrazole 14. The 2-methanesulfone group in 14 is conveniently displaced with various amines, aryloxides or alkoxides in solvents such ая tetrahydrofuran, dioxane, dimethylformamide acetonitrile at temperatures ranging from 20 °C to 200 °C. Under these reaction conditions the tosyl protecting group on the pyrazole is also simultaneously deprotected. Aqueous workup affords the desired tosyl deprotected, 2alkoxy, or 2-aryloxy or 2-amino substituted pyrazoles 15. The alkoxides or aryloxides are generated from their respective alcohols or phenols with suitable bases such as LiHMDS. NaH, LDA or tBuOK in solvents such tetrahydrofuran, dioxane ordimethylformamide. Deprotection of the remaining N-Boc group in 15 accomplished with trifluoroacetic acid or hydrochloric acid in solvents such as dichloromethane or dioxane to afford the pyrazole 16. Treatment of the pyrazole 16 with an acid chloride 7 in the presence of base or with an acid 8 under standard peptide coupling conditions (EDC or DCC or PyBrOP with an additive such as HOBt or HATU and base

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such as N-methylmorpholine or diisopropyl ethylamine) affords the desired final products 17.

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The following 2-substituted pyrimidine compounds can be prepared as set forth above, particularly in a manner similar to that outlined above in Scheme D-2.

5 Example D-34

5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-thiomethyl)pyrimidyl]-3-(4-chlorophenyl)pyrazole

Example D-35

5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-methanesulfonyl)pyrimidyl]-3-(4-chlorophenyl)pyrazole

Example D-36

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5-[4-N-(2-hydroxyacety1)piperidy1]-4-[4-(2-amino)pyrimidy1]-3-(4-chloropheny1)pyrazole

Example D-37

5 5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-methylamino)pyrimidyl]-3-(4-chlorophenyl)pyrazole

10 Example D-38

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5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-isopropylamino)pyrimidyl]-3-(4-chlorophenyl)pyrazole

Example D-39

5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-S-methylbenzylamino)pyrimidyl]-3-(4-chlorophenyl)pyrazole

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Example D-40

5 5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(2-R-methylbenzylamino)pyrimidyl]-3-(4-chlorophenyl)pyrazole

10 Example D-41

5-[4-N-(2-hydroxyacety1)piperidy1]-4-[4-(2-methoxy)pyrimidy1]-3-(4-chloropheny1)pyrazole

Example D-42

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5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(p-fluorophenoxy)pyrimidyl]-3-(4-chlorophenyl)pyrazole

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5-[4-N-(2-hydroxyacetyl)piperidyl]-4-[4-(p-fluoroanilino)pyrimidyl]-3-(4-chlorophenyl)pyrazole

Example D-43

CI N NH

In a manner similar to that outlined above in Scheme D-1, for the synthesis of the piperidine analogs 6, the aminocyclohexane analogs are prepared by substitution of 1 in Scheme D-1 with a suitably protected (Boc is shown) methyl or ethyl ester of cis-aminocyclohexane carboxylic acid 10 or trans-aminocyclohexane carboxylic acid 11 or trans-aminomethylcyclohexane carboxylic acid 12, which

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affords the cis-aminocyclohexane 13, or transaminocyclohexane 14 or the trans-aminomethylcyclohexane 15
respectively (Scheme 3). Suitable reductive alkylations
on 13, 14 or 15 with 1-1.5 equivalents of aldehydes or
ketones in the presence of a reducing agent like sodium
cyanoborohydride or sodium triacetoxyborohydride in
solvents such as methanol, ethanol, acetic acid,
tetrahydrofuran or dichloromethane lead to the desired
mono-alkylated derivatives 16, 17 or 18 respectively.

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Scheme 3

where R4 can be H

The dimethyl derivatives 19, 20 or 21 can be prepared by heating a solution of the aminocyclohexanes 13, 14 or 15

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respectively in a mixture of formaldehyde and formic acid at temperatures ranging from 40 °C to 110 °C.

An additional group of compounds of interest includes 10 the following:

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Biological data for a number of compounds are shown in the following table. In vitro p38 alpha kinase inhibitory data are shown in the column identified as "p38 5 alpha IC_{so} (μM)". In vitro human whole blood assay data for measuring the ability of the compounds to inhibit TNF production in human whole blood stimulated with LPS are shown in the column identified as: "HWB IC_{50} (μ M)". vivo assessment of the ability of the compounds to inhibit LPS-stimulated TNF-release in the rat is shown in the column identified as: "ratLPS/%Inh@dose(mg/kg)" wherein the dose is in milligram per kilogram (mg/kg) administered by oral gavage, 4 hours before LPS challenge.

Example	p38 alpha	HWB IC ₅₀	ratLPS/%Inh	ratLPS/%Inh	ratLPS/%Inh
	IC ₅₀ (uM)	(uM)	@1.0(mg/kg)	@5.0(mg/kg)	@20.0(mg/kg)
D-1	0.17		83.0		
D-2	0.084	1.79	89.0	95.0	
D-3	0.095	0.46	69.0	88.0	91.0
D-4	0.91	1.55	42.3	83.0	99.0
D-5	0.14	4.09	65.0	78.5	83.0
D-6	0.083	1.33	82.0	96.0	100
D-7	0.44	>25.0		0	
D-8	0.18	1.3	65	85	
D-9	1.63	15.8	5	86	
D-10	3.95	14.8		80	
D-11	0.16	1.5	43	86	
D-12	0.82	7.06	71	91	
D-13	0.33	8.36	53	87	-

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WHAT IS CLAIMED IS:

1. A compound of Formula IB:

wherein

R1 is selected from hydrido, hydroxy, alkyl, 5 cycloalkyl, alkenyl, cycloalkenyl, alkynyl, arvl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene. heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, 10 hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, 15 alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene, 20 alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, 25 alkylcarbonylalkylene, arylcarbonylalkylene,

alkylcarbonylalkylene, arylcarbonylalkylene,
heterocyclylcarbonylalkylene, alkylcarbonylarylene,
arylcarbonylarylene, heterocyclylcarbonylarylene,
alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene,
heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,

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30 arylcarbonyloxyarylene, and heterocyclylcarbonyloxyarylene; or

R1 has the formula

wherein:

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i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, and

heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl,

- aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclylalkylene, alkylheterocyclylarylene,
- aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,
- alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene,
- aryloxycarbonylarylene, alkylaryloxycarbonylarylene, arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene,

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alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, cycloalkylthioalkylene, alkylthioarylene, 65 aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, 70 alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthicalklylarylene, and alkylsulfonylarylene groups 75 may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or

R²⁷ is -CHR²⁸R²⁹ wherein R²⁸ is alkoxycarbonyl, and R²⁹ is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl and nitro; or

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R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy; and

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R2 is piperidinyl substituted with one or more 100 substituents selected from hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, alkoxyalkylene, alkoxyalkenylene, alkoxyalkynylene, and hydroxyacyl, wherein said hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, alkoxyalkylene, alkoxyalkenylene, alkoxyalkynylene, and 105 hydroxyacyl substitutents may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be 110 optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or 115 R2 is piperidinyl substituted with one or more substituents selected from hydroxycycloalkyl and alkoxycycloalkyl, and wherein said hydroxycycloalkyl and alkoxycycloalkyl substitutents may be optionally substituted with one or more substituents selected from 120 cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, 125 alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and R3 is selected from pyridinyl, pyrimidinyl,

quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

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groups may be optionally substituted with one or more substituents independently selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R4 is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R4 is optionally substituted with one or more substituents independently selected from halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer

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thereof.

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2. A compound of Claim 1 wherein:

R² is piperidinyl substituted with one or more substituents selected from hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, alkoxyalkylene, alkoxyalkenylene, alkoxyalkynylene, hydroxyalkylcarbonyl, hydroxyalkenylcarbonyl, and hydroxyalkynylcarbonyl, wherein said hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, alkoxyalkylene, alkoxyalkenylene, alkoxyalkynylene, hydroxyalkylcarbonyl, hydroxyalkenylcarbonyl, and hydroxyalkynylcarbonyl substitutents may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, alkyl, aryl, arylalkyl, haloalkyl, aryl, arylalkyl, haloalkyl, and

heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

R² is piperidinyl substituted with one or more substituents selected from hydroxycycloalkyl, alkoxycycloalkyl, and hydroxycycloalkyl, and hydroxycycloalkyl, and hydroxycycloalkylcarbonyl substitutents may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, baloalkyl, and heteroarylalkyl, baloalkyl, and heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy.

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3. A compound of Claim 1 selected from compounds, their tautomers and their pharmaceutically acceptable salts, of the group consisting of:

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4. A compound of Claim 1 having Formula XB:

wherein

Z represents a carbon atom or a nitrogen atom;

R1 is selected from hydrido, hydroxy, alkyl,

5 cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl,

10 arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl,

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alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino, 15 alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene, 20 alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene. heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, arylcarbonylalkylene, 25 heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene, arylcarbonyloxyarylene, and 30 heterocyclylcarbonyloxyarylene; and ${\ensuremath{\mathbb{R}}}^2$ is piperidinyl substituted with one or more substituents selected from hydroxyalkyl, hydroxyalkenyl, alkoxyalkylene, alkoxyalkenylene, hydroxyalkylcarbonyl, and hydroxyalkenylcarbonyl, wherein said hydroxyalkyl, 35 hydroxyalkenyl, alkoxyalkylene, alkoxyalkenylene, hydroxyalkylcarbonyl, and hydroxyalkenylcarbonyl substitutents may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said 40 cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be optionally

keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl,
alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and
heteroaralkoxy; or

substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy,

R² is piperidinyl substituted with one or more substituents selected from hydroxycycloalkyl and hydroxycycloalkylcarbonyl, wherein said hydroxycycloalkyl and hydroxycycloalkylcarbonyl substitutents may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be optionally substituted with one or more substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R4 is selected from cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R4 is optionally substituted with one or more substituents independently selected from halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R⁵ represents one or more substituents independently selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer

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thereof.

- 5. A compound of Claim 4 wherein R² is piperidinyl substituted with at least one substituent attached to the distal nitrogen heteroatom or to a carbon ring atom adjacent to the distal nitrogen heteroatom of the piperidine ring.
- 6. A compound of Claim 4 wherein Z represents a carbon atom.
- 7. A compound of Claim 4 wherein ${\bf Z}$ represents a nitrogen atom.
- 8. A compound of Claim 4 wherein R¹ is selected from hydrido, alkyl, hydroxyalkyl and alkynyl.
 - 9. A compound of Claim 4 wherein R1 is hydrido.
- 10. A compound of Claim 4 wherein R² is piperidinyl substituted with at least one substituent selected from lower hydroxyalkyl, lower hydroxyalkylcarbonyl and hydroxycycloalkylcarbonyl.
- 11. A compound of Claim 4 wherein R4 is optionally substituted phenyl.
- 12. A compound of Claim 4 wherein R' is phenyl optionally substituted at a substitutable position with one or more radicals independently selected from chloro, fluoro, bromo and iodo.
- 13. A compound of Claim 4 wherein R⁴ is phenyl optionally substituted at the meta or para position with one or more chloro radicals.

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- 14. A compound of Claim 4 wherein R⁵ is hydrido.
- 15. A compound of Claim 1 having Formula XX:

wherein:

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Z represents a carbon atom or a nitrogen atom; R400 is selected from hydroxyalkyl, 5 hydroxyalkylcarbonyl and alkoxyalkylene, wherein said hydroxyalkyl, hydroxyalkylcarbonyl and alkoxyalkylene may be optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl, wherein said cycloalkyl, 10 alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, 15 cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

R⁴⁰⁰ is hydroxycycloalkylcarbonyl that is optionally substituted with one or more substituents selected from cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and

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heteroarylalkyl, wherein said cycloalkyl, alkyl, aryl, arylalkyl, haloalkyl, and heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from alkylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

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R^{401a} and R^{401b} are independently selected from hydrogen, halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R⁴⁰² is selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

16. A compound of Claim 15 wherein:

R⁴⁰⁰ is selected from lower hydroxyalkyl, lower hydroxyalkylcarbonyl and lower alkoxyalkylene, wherein said lower hydroxyalkyl, lower hydroxyalkylcarbonyl and lower alkoxyalkylene may be optionally substituted with one or more substituents selected from cycloalkyl, lower alkyl, phenyl, lower phenylalkyl, lower haloalkyl, and

lower heteroarylalkyl, wherein said cycloalkyl, lower alkyl, phenyl, lower phenylalkyl, lower haloalkyl, and lower heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from lower alkylene, lower alkynylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; or

R400 is hydroxycycloalkylcarbonyl that is optionally substituted with one or more substituents selected from cycloalkyl, lower alkyl, phenyl, lower phenylalkyl, lower haloalkyl, and lower heteroarylalkyl, wherein said cycloalkyl, lower alkyl, phenyl, lower phenylalkyl, lower haloalkyl, and lower heteroarylalkyl substituents may be optionally substituted with one or more substituents selected from lower alkylene, lower alkynylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, aryloxy, heterocyclyl, and lower heteroaralkoxy; and

R^{401a} and R^{401b} are independently selected from hydrogen, halo, lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl, wherein said lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl substituents may be optionally substituted with one or more lower alkylene, lower alkenylene, lower alkynylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; and

R⁴⁰² is selected from hydrogen, phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy, wherein said

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phenyl, lower alkylamino, lower alkylthio, lower
alkyloxy, phenyloxy, phenylamino, phenylthio, and
45 phenylalkoxy may be optionally substituted with one or
more lower alkylene, lower alkenylene, hydroxy, halo,
lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano,
lower alkylsulfonyl, lower alkylsulfinyl, lower
alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl,
50 and lower heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

- 17. A compound of Claim 15 wherein Z represents a carbon atom.
- 18. A compound of Claim 15 wherein Z represents a nitrogen atom.
- 19. A compound of Claim 15 wherein R^{400} is optionally substituted hydroxyalkylcarbonyl.
- 20. A compound of Claim 15 wherein \mathbb{R}^{400} is optionally substituted hydroxycycloalkylcarbonyl.
- 21. A compound of Claim 15 wherein R⁴⁰⁰ is optionally substituted alkoxyalkylene.
- 22. A compound of Claim 15 wherein R⁴⁰⁰ is optionally substituted hydroxyalkyl.
- 23. A compound of Claim 15 wherein \mathbb{R}^{401} represents one or more chloro, fluoro, bromo and iodo.
- 24. A compound of Claim 15 wherein \mathbb{R}^{401} is metachloro or para-chloro.
 - 25. A compound of Claim 15 wherein R402 is hydrido.

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26. A compound of Claim 15 wherein:

R⁴⁰⁰ is optionally substituted lower hydroxyalkylcarbonyl;

 ${\rm R}^{\rm 401a}$ is selected from chloro, fluoro, bromo and iodo; and

R402 is hydrido.

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27. A compound of Claim 15 wherein:

R⁴⁰⁰ is selected from optionally substituted 2-hydroxyacetyl, 2-hydroxy-proprionyl, 2-hydroxy-2-methylpropionyl, 2-hydroxy-2-phenylacetyl, 3-

hydroxyproprionyl, 2-hydroxy-3-methylbutyryl, 2hydroxyisocapropyl, 2-hydroxy-3-phenylproprionyl, and 2hydroxy-3-imidazolylproprionyl;

 ${\rm R}^{\rm 401a}$ is selected from chloro, fluoro, bromo and iodo; and

10 R402 is hydrido.

- 28. A compound of Claim 27 wherein \mathbb{R}^{401a} is metachloro or para-chloro.
- 29. A compound of Claim 27 wherein R^{401a} is parachloro and R^{401b} is hydrogen.
 - 30. A compound of Claim 15 wherein:

R⁴⁰⁰ is optionally substituted lower hydroxycycloalkylcarbonyl;

 \mathbf{R}^{401a} is selected from chloro, fluoro, bromo and iodo; and

R402 is hydrido.

31. A compound of Claim 15 wherein:

 ${\tt R^{400}}$ is selected from optionally substituted 1-hydroxy-1-cyclohexylacetyl, 2-hydroxy-1-cyclohexylacetyl, 4-hydroxy-1-

5 cyclohexylacetyl, 1-hydroxy-1-cyclopentylacetyl, 2-

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hydroxy-1-cyclopentylacetyl, and 3-hydroxy-1-cyclopentylacetyl, 2-hydroxy-2-cyclohexylacetyl;

 R^{401a} is selected from chloro, fluoro, bromo and iodo; and

10 R⁴⁰² is hydrido.

- 32. A compound of Claim 31 wherein \mathbb{R}^{401a} is metachloro or para-chloro.
 - 33. A compound of Claim 15 wherein:

 R^{400} is optionally substituted lower hydroxyalkyl;

R401 is selected from chloro, fluoro, bromo and iodo;

and

5 R⁴⁰² is hydrido.

34. A compound of Claim 15 wherein:

R⁴⁰⁰ is selected from optionally substituted hydroxymethyl, hydroxyethyl, hydroxypropyl and hydroxyisopropyl;

5 R^{401a} is selected from chloro, fluoro, bromo and iodo; and

R402 is hydrido.

- 35. A compound of Claim 34 wherein R^{401a} is metachloro or para-chloro.
 - 36. A compound of Claim 15 wherein:

 R^{400} is optionally substituted lower alkoxyalkylene; R^{401a} is selected from chloro, fluoro, bromo and iodo;

R402 is hydrido.

and

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37. A compound of Claim 15 wherein:

R⁴⁰⁰ is selected from optionally substituted methoxymethylene, methoxyethylene, methoxypropylene, methoxyisopropylene, ethoxymethylene, ethoxyethylene,

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5 ethoxypropylene, and ethoxyisopropylene.

 ${\rm R}^{\rm 401a}$ is selected from chloro, fluoro, bromo and iodo; and

R402 is hydrido.

38. A compound of Claim 37 wherein R^{401a} is metachloro or para-chloro.

39. A compound of Formula IC:

wherein

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R¹ is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl,

- haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl,
- alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl,
- 20 alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl,
 heterocyclylsulfonyl, alkylaminoalkylene,
 alkylsulfonylalkylene, acyl, acyloxycarbonyl,
 alkoxycarbonylalkylene, aryloxycarbonylalkylene,

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heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene,
aryloxycarbonylarylene, heterocyclyloxycarbonylarylene,
alkylcarbonylalkylene, arylcarbonylalkylene,
heterocyclylcarbonylalkylene, alkylcarbonylarylene,
arylcarbonylarylene, heterocyclylcarbonylarylene,
alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene,
heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,
arylcarbonyloxyarylene, and
heterocyclylcarbonyloxyarylene; or

R1 has the formula

35 wherein:

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i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl,

alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclyl, alkylheterocyclylalkylene, alkylheterocyclylarylene,

aralkylneterocyclylalkylene, alkylneterocyclylarylene, aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene,

alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl, alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl,

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alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, 60 arvlcarbonvlalkylene, alkoxycarbonvlarvlene, aryloxycarbonylarylene, alkylaryloxycarbonylarylene, arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, 65 heterocyclylcarbonylalkylarylene, alkylthioalkylene, cycloalkylthioalkylene, alkylthioarylene, aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein 70 said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, 75 arylthioalklylarylene, and alkylsulfonylarylene groups may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or

R²⁷ is -CHR²⁶R²⁹ wherein R²⁸ is alkoxycarbonyl, and R²⁹
is selected from aralkyl, aralkoxyalkylene,
heterocyclylalkylene, alkylheterocyclylalkylene,
alkoxycarbonylalkylene, alkylthioalkylene, and
aralkylthioalkylene; wherein said aralkyl and
heterocylcyl groups may be optionally substituted with
one or more radicals independently selected from alkyl
and nitro; or

 R^{26} and R^{27} together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene,

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alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy; and

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R² is cyclohexyl substituted with one or more substituents selected from optionally substituted hydroxyalkyl, alkylaminoalkylene and cycloalkylamino; and

R³ is selected from pyridinyl, pyrimidinyl,
quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl,
thiazolylalkyl, thiazolylamino,

wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

groups may be optionally substituted with one or more substituents independently selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

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R' is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein

R' is optionally substituted with one or more substituents independently selected from halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

40. A compound of Claim 39 selected from compounds, their tautomers and their pharmaceutically acceptable salts, of the group consisting of :

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41. A compound of Claim 39 having Formula XC:

wherein

Z represents a carbon atom or a nitrogen atom; R¹ is selected from hydrido, hydroxy, alkyl,

- 5 cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl,
- arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino,
- 15 alkenylamino, alkynylamino, arylamino, heterocyclylamino,

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alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene,

- alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, arylcarbonylalkylene,
- heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene, arylcarbonyloxyarylene, and
- 30 heterocyclylcarbonyloxyarylene; and

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 $\ensuremath{R^2}$ is cyclohexyl substituted with one or more substituents selected from optionally substituted hydroxyalkyl, alkylaminoalkylene and cycloalkylamino; and

R⁴ is selected from cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R⁴ is optionally substituted with one or more substituents independently selected from halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R⁵ represents one or more substituents independently selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto,

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amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

55 a pharmaceutically-acceptable salt or tautomer thereof.

- 42. A compound of Claim 41 wherein R^2 is cyclohexyl substituted with at least one substituent attached to the 4-position carbon ring atom of the cyclohexyl ring.
- 43. A compound of Claim 41 wherein Z represents a carbon atom.
- 44. A compound of Claim 41 wherein Z represents a nitrogen atom.
- 45. A compound of Claim 41 wherein R¹ is selected from hydrido, alkyl, hydroxyalkyl and alkynyl.
 - 46. A compound of Claim 41 wherein R1 is hydrido.
- 47. A compound of Claim 41 wherein R^2 is cyclohexyl substituted with one or more substituents selected from optionally substituted lower hydroxyalkyl, lower alkylaminoalkylene and cycloalkylamino.
- 48. A compound of Claim 41 wherein R^4 is optionally substituted phenyl.
- 49. A compound of Claim 41 wherein R⁴ is phenyl optionally substituted at a substitutable position with one or more radicals independently selected from chloro, fluoro, bromo and iodo.
- 50. A compound of Claim 41 wherein R4 is phenyl optionally substituted at the meta or para position with

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one or more chloro radicals.

- 51. A compound of Claim 41 wherein R5 is hydrido.
- 52. A compound of Claim 41 having Formula XXIA:

wherein:

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and

Z represents a carbon atom or a nitrogen atom; R^{403} is selected from hydroxyalkyl,

alkylaminoalkylene and cycloalkylamino; and

R^{404a} and R^{404b} are independently selected from
hydrogen, halo, haloalkyl, haloalkoxy, alkoxy, cyano,
hydroxy, alkyl, alkenyl, and alkynyl, wherein said
haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl,
and alkynyl substituents may be optionally substituted
with one or more alkylene, alkenylene, alkynylene,
hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro,
cyano, alkylsulfonyl, alkylsulfinyl, alkylthio,
alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy;

R⁴⁰⁵ is selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy

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substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

53. A compound of Claim 52 wherein:

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 ${\tt R}^{403}$ is selected from lower hydroxyalkyl, lower alkylaminoalkylene and cycloalkylamino; and

R^{404a} and R^{404b} are independently selected from hydrogen, halo, lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl, wherein said lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl substituents may be optionally substituted with one or more lower alkylene, lower alkenylene, lower alkynylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; and

R⁴⁰⁵ is selected from hydrogen, phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy, wherein said phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy may be optionally substituted with one or more lower alkylene, lower alkenylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

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- 54. A compound of Claim 52 wherein ${\bf Z}$ represents a carbon atom.
- 55. A compound of Claim 52 wherein Z represents a nitrogen atom.
- 56. A compound of Claim 52 wherein R^{403} is optionally substituted hydroxyalkyl.
- 57. A compound of Claim 52 wherein R^{403} is optionally substituted alkylaminoalkylene.
- 58. A compound of Claim 57 wherein R^{403} is optionally substituted dialkylaminoalkylene.
- 59. A compound of Claim 52 wherein R403 is optionally substituted cycloalkylamino.
- 60. A compound of Claim 52 wherein R^{404a} is selected from chloro, fluoro, bromo and iodo.
- 61. A compound of Claim 52 wherein R^{404a} is metachloro or para-chloro.
 - 62. A compound of Claim 52 wherein R^{405} is hydrido.
 - 63. A compound of Claim 52 wherein:

 R^{403} is optionally substituted lower hydroxyalkyl; R^{404a} is selected from chloro, fluoro, bromo and iodo;

5 and

R405 is hydrido.

64. A compound of Claim 52 wherein:

R⁴⁰³ is selected from optionally substituted hydroxymethyl, hydroxyethyl, hydroxypropyl and hydroxybutyl;

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5 R^{404a} is selected from chloro, fluoro, bromo and iodo; and

R405 is hydrido.

- 65. A compound of Claim 64 wherein R^{404a} is metachloro or para-chloro.
 - 66. A compound of Claim 52 wherein:

R⁴⁰³ is optionally substituted lower alkylaminoalkylene;

R404a is selected from chloro, fluoro, bromo and iodo;

5 and

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and

R405 is hydrido.

67. A compound of Claim 52 wherein:

R⁴⁰³ is selected from optionally substituted methylaminomethylene, methylaminoethylene, methylaminopropylene, ethylaminomethylene, ethylaminoethylene, ethylaminoethylene, propylaminomethylene, propylaminomethylene, propylaminopropylene, dimethylaminomethylene, dimethylaminopropylene, diethylaminomethylene, diethylaminomethylene, diethylaminopropylene, diethylaminopropylene, dipropylaminomethylene, dipropylaminomethylene, dipropylaminopropylene;

 $R^{404\hat{a}}$ is selected from chloro, fluoro, bromo and iodo; and

R405 is hydrido.

- 68. A compound of Claim 67 wherein R^{404a} is metachloro or para-chloro.
 - 69. A compound of Claim 52 wherein: $R^{403} \mbox{ is optionally substituted cycloalkylamino;} \\ R^{404a} \mbox{ is selected from chloro, fluoro, bromo and iodo;}$

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5 R405 is hydrido.

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70. A compound of Claim 52 wherein: $R^{403} \text{ is selected from optionally substituted} \\$ cyclopropyl, cyclobutyl, cyclopentyl and cyclohexyl; $R^{404a} \text{ is selected from chloro, fluoro, bromo and iodo;} \\$ and $R^{405} \text{ is hydrido.}$

71. A compound of Formula XXIB:

wherein:

Z represents a carbon atom or a nitrogen atom; R⁴⁰³ is selected from alkylamino; and R^{404a} and R^{404b} are independently selected from hydrogen, halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro,

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cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R405 is selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or 20 more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

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a pharmaceutically-acceptable salt or tautomer 25 thereof.

A compound of Claim 71 wherein: R403 is selected from lower alkylamino; and R404a and R404b are independently selected from hydrogen, halo, lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl, wherein said lower haloalkyl, lower haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and lower alkynyl substituents may be optionally substituted with one or more lower alkylene, lower alkenylene, lower alkynylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; and

R405 is selected from hydrogen, phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy, wherein said phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy may be optionally substituted with one or more lower alkylene, lower alkenylene, hydroxy, halo,

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lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

- 73. A compound of Claim 71 wherein Z represents a carbon atom.
- 74. A compound of Claim 71 wherein Z represents a nitrogen atom.
- 75. A compound of Claim 71 wherein R⁴⁰³ is optionally substituted dialkylamino.
- 76. A compound of Claim 71 wherein R^{404a} is selected from chloro, fluoro, bromo and iodo.
- 77. A compound of Claim 71 wherein \mathbb{R}^{4048} is metachloro or para-chloro.
 - 78. A compound of Claim 71 wherein R405 is hydrido.
- 79. A compound of Claim 71 wherein: $R^{403} \mbox{ is optionally substituted lower alkylamino;} \\ R^{404a} \mbox{ is selected from chloro, fluoro, bromo and iodo;} \\ \mbox{and}$
- 5 R⁴⁰⁵ is hydrido.

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80. A compound of Claim 71 wherein:

R⁴⁰³ is selected from optionally substituted
methylamino, ethylamino, n-propylamino, isopropylamino,
n-butylamino, sec-butylamino, t-butylamino,
isobutylamino, dimethylamino, diethylamino, di-npropylamino, di-isopropylamino, di-n-butylamino, di-sec-

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butylamino, di-t-butylamino, and di-isobutylamino; $R^{404a} \ \text{is selected from chloro, fluoro, bromo and iodo;}$ and

10 R⁴⁰⁵ is hydrido.

- 81. A compound of Claim 80 wherein \mathbb{R}^{404a} is metachloro or para-chloro.
 - 82. A compound Formula XXII:

wherein:

Z represents a carbon atom or a nitrogen atom; R^{406} is alkynyl; and

R407a and R407b are independently selected from hydrogen, halo, haloalkyl, haloalkoxy, alkoxy, cyano, hydroxy, alkyl, alkenyl, and alkynyl, wherein said haloalkyl, haloalkoxy, alkoxy, hydroxy, alkyl, alkenyl, and alkynyl substituents may be optionally substituted with one or more alkylene, alkenylene, alkynylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; and

R408 is selected from hydrogen, aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio,

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aralkoxy, wherein said aryl, alkylamino, alkylthio, alkyloxy, aryloxy, arylamino, arylthio, aralkoxy substituents may be optionally substituted with one or more alkylene, alkenylene, hydroxy, halo, haloalkyl, alkoxy, keto, amino, nitro, cyano, alkylsulfonyl, alkylsulfinyl, alkylthio, alkoxyalkyl, aryloxy, heterocyclyl, and heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer thereof.

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83. A compound of Claim 82 wherein:

R⁴⁰⁶ is selected from lower alkynyl; and
R^{407a} and R^{407b} are independently selected from
hydrogen, halo, lower haloalkyl, lower haloalkoxy, lower
alkoxy, cyano, hydroxy, lower alkyl, lower alkenyl, and
lower alkynyl, wherein said lower haloalkyl, lower
haloalkoxy, lower alkoxy, cyano, hydroxy, lower alkyl,
lower alkenyl, and lower alkynyl substituents may be
optionally substituted with one or more lower alkylene,
lower alkenylene, lower alkynylene, hydroxy, halo, lower
haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower
alkylsulfonyl, lower alkylsulfinyl, lower alkylthio,
lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower
heteroaralkoxy; and

R⁴⁰⁸ is selected from hydrogen, phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy, wherein said phenyl, lower alkylamino, lower alkylthio, lower alkyloxy, phenyloxy, phenylamino, phenylthio, and phenylalkoxy may be optionally substituted with one or more lower alkylene, lower alkenylene, hydroxy, halo, lower haloalkyl, lower alkoxy, keto, amino, nitro, cyano, lower alkylsulfonyl, lower alkylsulfinyl, lower alkylthio, lower alkoxyalkyl, phenyloxy, heterocyclyl, and lower heteroaralkoxy; or

a pharmaceutically-acceptable salt or tautomer

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thereof.

- $84.\ \ \mbox{A}$ compound of Claim 82 wherein Z represents a carbon atom.
- 85. A compound of Claim 82 wherein Z represents a nitrogen atom.
- 86. A compound of Claim 82 wherein \mathbb{R}^{407a} is selected from chloro, fluoro, bromo and iodo.
- 87. A compound of Claim 82 wherein R^{407a} is metachloro or para-chloro.
 - 88. A compound of Claim 82 wherein R408 is hydrido.
 - 89. A compound of Claim 82 wherein:

R406 is optionally substituted lower alkynyl;

 $\ensuremath{\text{R}}^{407a}$ is selected from chloro, fluoro, bromo and iodo; and

5 R408 is hydrido.

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90. A compound of Claim 82 wherein:

 \mathbb{R}^{406} is selected from optionally substituted ethynyl, propynyl and butynyl;

 R^{407a} is selected from chloro, fluoro, bromo and iodo; and

R408 is hydrido.

- 91. A compound of Claim 82 wherein R^{405} is propargyl.
- 92. A compound of Claim 82 wherein R^{407a} is metachloro or para-chloro.
 - 93. A compound of Formula IA

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$$R^{3}$$

$$R^{4}$$

$$\downarrow_{5}$$

$$\downarrow_{1}$$

$$\downarrow_{1$$

wherein

R1 is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, 5 heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, 10 arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, 15 alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene, 20 alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, arylcarbonylalkylene, 25 heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,

arylcarbonyloxyarylene, and

heterocyclylcarbonyloxyarylene; or

R¹ has the formula

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wherein:

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i is an integer from 0 to 9;

35 R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclyl,

alkylheterocyclylalkylene, alkylheterocyclylarylene, aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene,

alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl,

alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene,

aryloxycarbonylarylene, alkylaryloxycarbonylarylene,

arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, cycloalkylthioalkylene, alkylthioarylene,

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aralkylthioarylene, heterocyclylthioarylene, 65 arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, and alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, 70 alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups may be optionally substituted with one or more radicals 75 independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or

 R^{27} is $-CHR^{28}R^{29}$ wherein R^{28} is alkoxycarbonyl, and R^{29} is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl and nitro; or

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100

and

 R^{26} and R^{27} together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene,

alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be 95 optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy;

R² is selected from mercapto, aryl(hydroxyalkyl)amino, N-alkyl-N-alkynyl-amino, aminocarbonylalkylene, alkylcarbonylaminoalkylene,

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aminoalkylcarbonylaminoalkylene,
        alkylaminoalkylcarbonylamino, aminoalkylthio,
        alkylaminocarbonylalkylthio,
        alkylaminoalkylaminocarbonylalkylthio, cyanoalkylthio,
        alkenylthio, alkynylthio, carboxyalkylthio,
105
        alkoxycarbonylalkylthio, alkylsulfinyl, alkylsulfonyl,
        alkoxyalkyl, alkoxyalkylthio, alkoxycarbonylalkylamino,
        alkoxycarbonylaminoalkylene, alkoxycarbonylaminoalkoxy,
        aralkythio, heterocyclylalkylthio, aminoalkoxy,
110
        cyanoalkoxy, carboxyalkoxy, aryloxy, aralkoxy,
        alkenyloxy, alkynyloxy, and heterocyclylalkyloxy; or
              R^2 is R^{200}-heterocyclyl-R^{201}, R^{200}-aryl-R^{201}, or R^{200}-
        cycloalkyl-R201 wherein:
              R200 is selected from:
               - (CR202R203),-;
115
               -C(O)-:
               -C(O)-(CH2)-;
               -C(O)-O-(CH<sub>2</sub>),-;
               - (CH<sub>2</sub>),-C(O)-;
               -O-(CH<sub>2</sub>),-C(O)-;
120
               -NR<sup>202</sup>-;
               -NR^{202} - (CH_2)_{v} - ;
               - (CH<sub>2</sub>) ,-NR<sup>202</sup>-;
               - (CH<sub>2</sub>)<sub>y</sub>-NR<sup>202</sup>- (CH<sub>2</sub>)<sub>z</sub>-;
               - (CH<sub>2</sub>) , - C (O) - NR<sup>202</sup> - (CH<sub>2</sub>) , - ;
125
               -(CH_2)_v - NR^{202} - C(O) - (CH_2)_z - ;
               -(CH_2)_v - NR^{202} - C(O) - NR^{203} - (CH_2)_z - ;
               -S(O),-(CR202R203),-;
               - (CR202R203) y-S(O) x-;
               -S(O)_x-(CR^{202}R^{203})_y-O-;
130
               -S(O),-(CR202R203),-C(O)-;
               -O-(CH<sub>2</sub>),-;
               - (CH2),-O-;
               -S-; and
135
               -0-;
               or R200 represents a bond:
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R²⁰¹ represents one or more radicals selected from the group consisting of hydroxy, hydroxyalkyl, cycloalkyl, hydroxyalkylcarbonyl, cycloalkylcarbonyl,

- arylcarbonyl, haloarylcarbonyl, alkoxyalkylene,
 alkoxyarylene, carboxyalkylcarbonyl, alkoxyalkylcarbonyl,
 heterocyclylalkylcarbonyl, alkylsulfonylalkylene,
 aminoalkyl, aralkylamino, alkylaminoalkylene,
 aminocarbonyl, alkylcarbonylamino,
- alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl, alkylaminoalkylcarbonylamino, aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino, alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene, alkylimidocarbonyl, amidino, alkylamidino, aralkylamidino, guanidinoalkylene, and
 - alkylsulfonylamino; and R^{202} and R^{203} are independently selected from hydrido,

alkyl, aryl and aralkyl; and
 y and z are independently 0, 1, 2, 3, 4, 5 or 6
wherein y + z is less than or equal to 6; and

x is 0, 1 or 2; or

 R^2 is $\textsc{-NHCR}^{204}R^{205}$ wherein R^{204} is alkylaminoalkylene, and R^{205} is aryl; or

 R^2 is $-C(NR^{206})R^{207}$ wherein R^{206} is selected from hydrogen and hydroxy, and R^{207} is selected from alkyl, aryl and aralkyl; and

R³ is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylamino,

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155

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wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

groups may be optionally substituted with one or more

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200

radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkynylamino, cycloalkylamino, cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy,

aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylamino,

alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino, heterocyclylheterocyclylalkylamino,

heterocyclylheterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR⁴⁴R⁴⁵
wherein R⁴⁴ is alkylcarbonyl or amino, and R⁴⁵ is alkyl or aralkyl; and

R' is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R' is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl,

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alkylsulfinylalkylene, arylsulfinylalkylene, alkylsulfonyl, alkylsulfonylalkylene,

arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano, nitro, alkylamino, arylamino, alkylaminoalkylene, arylaminoalkylene, aminoalkylamino, and hydroxy;

provided R^3 is not 2-pyridinyl when R^4 is a phenyl ring containing a 2-hydroxy substituent and when R^1 is hydrido; and

further provided R^2 is selected from $-R^{200}$ -heterocyclyl- R^{201} , $-R^{200}$ -aryl- R^{201} , or $-R^{200}$ -unsubstituted cycloalkyl- R^{201} when R^4 is hydrido; and

further provided that R^4 is not methylsulfonylphenyl or aminosulfonylphenyl; and

further provided that R^{1} is not methylsulfonylphenyl; or

220 a pharmaceutically-acceptable salt or tautomer thereof.

94. A compound of Formula IXA:

wherein

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Z represents a carbon atom or a nitrogen atom; and \mathbb{R}^1 is selected from hydrido, lower alkyl, lower

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hydroxyalkyl, lower alkynyl, lower aralkyl, lower
     aminoalkyl and lower alkylaminoalkyl; and
           R<sup>2</sup> is lower hydroxyalkylamino; or
           R^2 is R^{200}-heterocyclyl-R^{201} or R^{200}-cycloalkyl-R^{201}
10
     wherein:
           R200 is selected from:
           - (CR202R203),-;
           -NR202-:
           -NR^{202} - (CH_2)_{v} - ;
           -(CH_2)_{v}-NR^{202}-;
15
           -O- (CH<sub>2</sub>),-;
           - (CH<sub>2</sub>) ,-O-;
           -S-;
           -0-;
           or R200 represents a bond;
20
          R^{201} represents one or more radicals selected from
     the group consisting of hydroxy, lower hydroxyalkyl,
     lower cycloalkyl, lower hydroxyalkylcarbonyl, lower
     cycloalkylcarbonyl, arylcarbonyl, haloarylcarbonyl, lower
25
     alkoxyalkylene, lower alkoxyarylene, lower
     carboxyalkylcarbonyl, lower alkoxyalkylcarbonyl, lower
     heterocyclylalkylcarbonyl, lower alkylsulfonylalkylene,
     amino, lower aminoalkyl, lower aralkylamino, lower
     alkylaminoalkylene, aminocarbonyl, lower
30
     alkylcarbonylamino, lower alkylcarbonylaminoalkylene,
     lower alkylaminoalkylcarbonyl, lower
     alkylaminoalkylcarbonylamino, lower
     aminoalkylcarbonylaminoalkyl, lower alkoxycarbonylamino,
     lower alkoxyalkylcarbonylamino, lower
35
     alkoxycarbonylaminoalkylene, lower alkylimidocarbonyl,
     amidino, lower alkylamidino, lower aralkylamidino,
     guanidino, lower guanidinoalkylene, and lower
     alkylsulfonylamino; and
          R^{202} and R^{203} are independently selected from hydrido,
40
     lower alkyl, aryl and lower aralkyl; and
          y is 0, 1, 2 or 3; and
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R4 is selected from aryl selected from phenyl, biphenyl, naphthyl, wherein said aryl is optionally substituted at a substitutable position with one or more radicals independently selected from halo, lower alkyl, lower alkoxy, aryloxy, lower aralkoxy, lower haloalkyl, lower alkylthio, lower alkylamino, nitro, and hydroxy; and

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R⁵ is selected from hydrido, halo, amino, cyano, aminocarbonyl, lower alkyl, lower alkoxy, hydroxy, lower aminoalkyl, lower aralkyl, lower aralkylamino, lower alkylamino, lower alkylamino, lower alkylamino, lower hydroxyalkylamino, lower alkylcarbonyl, lower aralkenyl, lower arylheterocyclyl, carboxy, lower cycloalkylamino, lower hydroxycycloalkylamino, lower alkoxycarbonylamino, lower alkoxyaralkylamino, lower alkylaminoalkylamino, lower heterocyclylamino, lower heterocyclylalkylamino, lower aralkylheterocyclylamino, lower alkylaminocarbonyl, lower alkylcarbonyl, lower alkylamino, hydrazinyl, and lower alkylhydrazinyl, or -NR⁶²R⁶³ wherein R⁶² is lower alkylcarbonyl or amino, and R⁶³ is lower alkyl or lower phenylalkyl; or

- a pharmaceutically-acceptable salt or tautomer $\,$ thereof.
 - 95. A compound of Claim 94 wherein R^2 is R^{200} -heterocyclyl- R^{201} .
 - 96. A compound of Claim 94 wherein R^2 is R^{200} -cycloalkyl- R^{201} .
 - 97. A compound of Claim 94 wherein:
 - R¹ is selected from hydrido, methyl, ethyl, hydroxyethyl and propargyl; and
 - $$R^2$$ is $R^{200}\mbox{-piperidinyl-}R^{201},~R^{200}\mbox{-piperazinyl-}R^{201},~or$ 5 $~R^{200}\mbox{-cyclohexyl-}R^{201}$ wherein:
 - R²⁰⁰ is selected from:

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- (CR202R203),-;
          -NR^{202}-:
          -S-:
10
          -0-;
          or R<sup>200</sup> represents a bond:
          R<sup>201</sup> represents one or more radicals selected from
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-
     1,1-dimethyl) ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
15
     cyclohexyl, methoxymethylene, methoxyethylene,
     methoxypropylene, ethoxyethylene, ethoxypropylene,
     propoxyethylene, propoxypropylene, methoxyphenylene,
     ethoxyphenylene, propoxyphenylene, cyclopropylcarbonyl,
20
     cyclobutylcarbonyl, cyclopentylcarbonyl,
     cyclohexylcarbonyl, benzoyl, chlorobenzoyl,
     fluorobenzoyl, hydroxymethylcarbonyl,
     hydroxyethylcarbonyl, hydroxypropylcarbonyl,
     carboxymethylcarbonyl, carboxyethylcarbonyl,
     carboxypropylcarbonyl, methoxymethylcarbonyl,
25
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl,
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
30
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
     methylsulfonvlmethylene, amino, aminomethyl, aminoethyl,
     aminopropyl, phenylamino, benzylamino,
35
     methylaminomethylene, ethylaminomethylene,
     methylaminoethylene, ethylaminoethylene, aminocarbonyl,
     methylcarbonylamino, ethylcarbonylamino,
     methylaminomethylcarbonyl, ethylaminomethylcarbonyl,
     methylcarbonylaminomethylene,
40
     ethylcarbonylaminomethylene,
     aminomethylcarbonylaminocarbonylmethylene,
     methoxycarbonylamino, ethoxycarbonylamino,
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methoxymethylcarbonylamino, methoxyethylcarbonylamino, ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,
45 methoxycarbonylaminomethylene, ethoxycarbonylaminomethylene, methylimidocarbonyl, ethylimidocarbonyl, amidino, methylamidino, methylamidino, benzylamidino, guanidino, guanidinomethylene, guanidinoethylene, and
50 methylsulfonylamino; and

 R^{202} and R^{203} are independently selected from hydrido, methyl, ethyl, propyl, butyl, phenyl and benzyl; and y is 0, 1 or 2; and

R4 is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from methylthio, fluoro, chloro, bromo, iodo, methyl, ethyl, methoxy, ethoxy, phenoxy, benzyloxy, trifluoromethyl, nitro, dimethylamino, and hydroxy; and

55

R⁵ is selected from hydrido, fluoro, chloro, bromo, iodo, hydroxy, methyl, ethyl, propyl, benzyl, fluorophenylethyl, fluorophenylethenyl, fluorophenylpyrazolyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, dimethylamino, 2-methylbutylamino, ethylamino,

dimethylaminoethylamino, hydroxyethylamino, hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino, hydroxycyclopentylamino, hydroxycyclohexylamino, imidazolylamino, morpholinylethylamino, (1-ethyl-2-

hydroxy) ethylamino, piperidinylamino, pyridinylmethylamino, phenylmethylpiperidinylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino,

fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, methylaminopropylamino, dimethylaminopropylamino, methylaminobutylamino, dimethylaminobutylamino, methylaminopentylamino,

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dimethylaminopentylamino, ethylaminoethylamino,
diethylaminoethylamino, ethylaminopropylamino,
diethylaminopropylamino, ethylaminobutylamino,
diethylaminobutylamino, ethylaminopentylamino,
methylaminocarbonyl, methylcarbonyl, ethylcarbonyl,
hydrazinyl, and 1-methylhydrazinyl, or -NR⁶²R⁶³ wherein R⁶²
is methylcarbonyl or amino, and R⁶³ is methyl or benzyl;
or

a pharmaceutically-acceptable salt or tautomer thereof.

- 98. A compound of Claim 97 wherein R^2 is R^{200} -piperidinyl- R^{201} .
- 99. A compound of Claim 97 wherein R^2 is R^{200} -pyrazinyl- R^{201} .
- 100. A compound of Claim 97 wherein $\ensuremath{R^2}$ is $\ensuremath{R^{200}}\xspace$ cyclohexyl- $\ensuremath{R^{201}}\xspace$.
 - 101. A compound of Claim 94 having the Formula XA:

wherein:

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Z represents a carbon atom or a nitrogen atom; and R¹ is selected from hydrido, methyl, ethyl,

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hydroxyethyl and propargyl; and
          R<sup>2</sup> is R<sup>200</sup>-piperidinyl-R<sup>201</sup> wherein:
          R<sup>200</sup> is selected from:
          - (CR202R203),-;
10
          -NR<sup>202</sup>-;
          -S-:
          -0-;
          or R<sup>200</sup> represents a bond;
          R<sup>201</sup> represents one or more radicals selected from
15
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-
     1,1-dimethyl) ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
     cyclohexyl, methoxymethylene, methoxyethylene,
     methoxypropylene, ethoxyethylene, ethoxypropylene,
20
     propoxyethylene, propoxypropylene, methoxyphenylene,
     ethoxyphenylene, propoxyphenylene, cyclopropylcarbonyl,
     cyclobutylcarbonyl, cyclopentylcarbonyl,
     cyclohexylcarbonyl, benzoyl, chlorobenzoyl,
     fluorobenzoyl, hydroxymethylcarbonyl,
25
     hydroxyethylcarbonyl, hydroxypropylcarbonyl,
     carboxymethylcarbonyl, carboxyethylcarbonyl,
     carboxypropylcarbonyl, methoxymethylcarbonyl,
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl,
30
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
35
     methylsulfonylmethylene, amino, aminomethyl, aminoethyl,
     aminopropyl, N-methylamino, N,N-dimethylamino, N-
     ethylamino, N, N-diethylamino, N-propylamino, N, N-
     dipropylamino, phenylamino, benzylamino,
     methylaminomethylene, ethylaminomethylene,
40
     methylaminoethylene, ethylaminoethylene, aminocarbonyl,
     methylcarbonylamino, ethylcarbonylamino,
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methylaminomethylcarbonyl, ethylaminomethylcarbonyl, methylcarbonylaminomethylene, ethylcarbonylaminomethylene,

- aminomethylcarbonylaminocarbonylmethylene,
 methoxycarbonylamino, ethoxycarbonylamino,
 methoxymethylcarbonylamino, methoxyethylcarbonylamino,
 ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,
 methoxycarbonylaminomethylene,
- ethoxycarbonylaminomethylene, methylimidocarbonyl, ethylimidocarbonyl, amidino, methylamidino, methylamidino, benzylamidino, guanidino, guanidinomethylene, guanidinoethylene, and methylsulfonylamino; and
 - R^{202} and R^{203} are independently selected from hydrido, methyl, ethyl, propyl, butyl, phenyl and benzyl; and

y is 0, 1 or 2; and

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R4 is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, ethyl, methoxy and ethoxy; and

R⁵ is selected from hydrido, fluoro, chloro, bromo, hydroxy, methyl, ethyl, propyl, benzyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, dimethylamino, 2-methylbutylamino,

- 65 methylamino, dimethylamino, 2-methylbutylamino, ethylamino, dimethylaminoethylamino, hydroxyethylamino, hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino,
- imidazolylamino, morpholinylethylamino, (1-ethyl-2-hydroxy)ethylamino, piperidinylamino, pyridinylmethylamino, phenylmethylpiperidinylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino,

hydroxycyclopentylamino, hydroxycyclohexylamino,

75 phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, methylaminopropylamino,

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dimethylaminopropylamino, methylaminobutylamino.
     dimethylaminobutylamino, methylaminopentylamino.
8.0
     dimethylaminopentylamino, ethylaminoethylamino,
     diethylaminoethylamino, ethylaminopropylamino,
     diethylaminopropylamino, ethylaminobutylamino,
     diethylaminobutylamino, ethylaminopentylamino,
     methylaminocarbonyl, methylcarbonyl, and ethylcarbonyl;
85
     or
          a pharmaceutically-acceptable salt or tautomer
     thereof.
                A compound of Claim 101 wherein:
          R1 is selected from hydrido, methyl, ethyl,
     hydroxyethyl and propargyl: and
          R2 is R200-piperidinyl-R201 wherein:
 5
          R200 is selected from:
          methylene;
          -NR<sup>202</sup>-:
          -S-;
          -0-;
          or R200 represents a bond;
10
          R^{201} represents one or more radicals selected from
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, (1-hydroxy-1,1-
     dimethyl) ethyl, methoxymethyl, methoxyethyl,
     methoxypropyl, ethoxyethyl, ethoxypropyl, propoxyethyl,
15
     propoxypropyl, methoxyphenyl, ethoxyphenyl,
     propoxyphenyl, hydroxymethylcarbonyl,
     hydroxyethylcarbonyl, carboxymethylcarbonyl,
     carboxyethylcarbonyl, methoxymethylcarbonyl,
20
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl.
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
25
    propoxyphenylcarbonyl, methylsulfonylmethylene, amino,
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aminomethyl, aminoethyl, aminopropyl, N-benzylamino, methylaminomethylene, aminocarbonyl, methoxycarbonylamino, ethoxycarbonylamino, or methylsulfonylamino; and

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 $\mbox{\ensuremath{R^{202}}}$ is selected from hydrido, methyl, ethyl, phenyl and benzyl; and

R⁴ is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, ethyl, methoxy and ethoxy; and

R⁵ is selected from hydrido, fluoro, chloro, bromo, hydroxy, methyl, ethyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, dimethylamino, ethylamino, dimethylamino, hydroxyethylamino, hydroxypropylamino, hydroxybutylamino,

hydroxyethylamino, hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino, hydroxycyclopentylamino, hydroxycyclohexylamino, (1-ethyl-2-hydroxy)ethylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino, phenylmethylamino.

methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, methylaminopropylamino, dimethylaminopropylamino, methylaminobutylamino, dimethylaminobutylamino, methylaminopentylamino, dimethylaminopentylamino

methylaminopentylamino, dimethylaminopentylamino, ethylaminoethylamino, diethylaminoethylamino, ethylaminopropylamino, diethylaminopropylamino, ethylaminobutylamino, diethylaminobutylamino, ethylaminopentylamino, methylaminocarbonyl, methylaminopentyl, and ethylaminopentyl; or

a pharmaceutically-acceptable salt or tautomer thereof.

103. A compound of Claim 101 wherein: R^1 is hydrido; and R^2 is R^{200} -piperidinyl- R^{201} wherein:

```
R200 is selected from:
 5
           methylene;
           -NR<sup>202</sup>-:
           -S-:
           -0-:
           or R<sup>200</sup> represents a bond;
10
          R^{201} represents one or more radicals selected from
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, methoxymethyl, methoxyethyl,
     methoxypropyl, ethoxyethyl, ethoxypropyl, propoxyethyl,
     propoxypropyl, methoxyphenyl, ethoxyphenyl,
15
     propoxyphenyl, hydroxymethylcarbonyl,
     hydroxyethylcarbonyl, carboxymethylcarbonyl,
     carboxyethylcarbonyl, methoxymethylcarbonyl,
     methoxyethylcarbonyl, ethoxymethylcarbonyl,
     ethoxyethylcarbonyl, methoxyphenylcarbonyl,
20
     ethoxyphenylcarbonyl, amino, aminomethyl, aminoethyl,
     aminopropyl, N-benzylamino, methylaminomethylene,
     aminocarbonyl, methoxycarbonylamino, and
     ethoxycarbonylamino; and
          R^{202} is selected from hydrido, methyl phenyl and
25
     benzyl; and
          R^4 is phenyl, wherein said phenyl is optionally
     substituted with one or more radicals independently
     selected from fluoro, chloro, methyl, and methoxy; and
          R<sup>5</sup> is selected from hydrido, methylamino,
     dimethylamino, 2-methylbutylamino, ethylamino,
30
     dimethylaminoethylamino, hydroxypropylamino,
     hydroxyethylamino, hydroxypropylamino, hydroxybutylamino,
     hydroxycyclopropylamino, hydroxycyclobutylamino,
     hydroxycyclopentylamino, hydroxycyclohexylamino, (1-
35
     ethyl-2-hydroxy) ethylamino, aminomethyl,
     cyclopropylamino, amino, dimethylaminoethylamino,
     dimethylaminopropylamino, dimethylaminobutylamino,
     dimethylaminopentylamino, diethylaminoethylamino,
     diethylaminopropylamino, diethylaminobutylamino, and
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40 diethylaminopentylamino; or

a pharmaceutically-acceptable salt or tautomer thereof.

104. A compound of Claim 101 wherein:

R1 is hydrido; and

R² is R²⁰⁰-piperidinyl-R²⁰¹ wherein:

R200 is selected from:

5 methylene;

15

20

-NR²⁰²-:

-S-:

-0~:

or R200 represents a bond;

10 R²⁰¹ represents one or more radicals selected from the group consisting of methoxyethyl, methylcarbonyl, hydroxymethylcarbonyl, methoxymethylcarbonyl, and amino; and

R²⁰² is selected from hydrido and methyl; and R⁴ is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, and methoxy; and

R⁵ is selected from hydrido, hydroxypropylamino, hydroxycyclohexylamino, diethylaminoethylamino; or

a pharmaceutically-acceptable salt or tautomer thereof. $% \left(1\right) =\left(1\right) \left(1$

105. A compound of Claim 94 having the Formula XA:

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wherein:
           Z represents a carbon atom or a nitrogen atom; and
 5
          R1 is selected from hydrido, methyl, ethyl,
     hydroxyethyl and propargyl; and
          R^2 is R^{200}-piperazinyl-R^{201} wherein:
          R<sup>200</sup> is selected from:
           - (CR<sup>202</sup>R<sup>203</sup>),-;
10
           -NR^{202}-;
          -S-:
          -0-:
          or R<sup>200</sup> represents a bond;
          R^{201} represents one or more radicals selected from
15
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-
     1,1-dimethyl)ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
     cyclohexyl, methoxymethylene, methoxyethylene,
     methoxypropylene, ethoxyethylene, ethoxypropylene,
20
     propoxyethylene, propoxypropylene, methoxyphenylene,
     ethoxyphenylene, propoxyphenylene, cyclopropylcarbonyl,
     cyclobutylcarbonyl, cyclopentylcarbonyl,
     cyclohexylcarbonyl, benzoyl, chlorobenzoyl,
     fluorobenzoyl, hydroxymethylcarbonyl,
25
     hydroxyethylcarbonyl, hydroxypropylcarbonyl.
     carboxymethylcarbonyl, carboxyethylcarbonyl,
     carboxypropylcarbonyl, methoxymethylcarbonyl,
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl,
30
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
     methylsulfonylmethylene, amino, aminomethyl, aminoethyl,
35
     aminopropyl, phenylamino, benzylamino,
     methylaminomethylene, ethylaminomethylene,
     methylaminoethylene, ethylaminoethylene, aminocarbonyl,
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1144

methylcarbonylamino, ethylcarbonylamino, 40 methylaminomethylcarbonyl, ethylaminomethylcarbonyl, methylcarbonylaminomethylene, ethylcarbonylaminomethylene, aminomethylcarbonylaminocarbonylmethylene, methoxycarbonylamino, ethoxycarbonylamino, 45 methoxymethylcarbonylamino, methoxyethylcarbonylamino, ethoxymethylcarbonylamino, ethoxyethylcarbonylamino, methoxycarbonylaminomethylene, ethoxycarbonylaminomethylene, methylimidocarbonyl, ethylimidocarbonyl, amidino, methylamidino, 50 methylamidino, benzylamidino, quanidino, guanidinomethylene, guanidinoethylene, and methylsulfonylamino; and R^{202} and R^{203} are independently selected from hydrido, methyl, ethyl, propyl, butyl, phenyl and benzyl; and 55 y is 0, 1 or 2; and R4 is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, ethyl, methoxy and ethoxy; and 60 R⁵ is selected from hydrido, fluoro, chloro, bromo, hydroxy, methyl, ethyl, propyl, benzyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, dimethylamino, 2-methylbutylamino, ethylamino, dimethylaminoethylamino, hydroxyethylamino, 65 hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino, hydroxycyclopentylamino, hydroxycyclohexylamino, imidazolylamino, morpholinylethylamino, (1-ethyl-2hydroxy) ethylamino, piperidinylamino, 70 pyridinylmethylamino, phenylmethylpiperidinylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino,

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75
     dimethylaminoethylamino, methylaminopropylamino,
     dimethylaminopropylamino, methylaminobutylamino,
     dimethylaminobutylamino, methylaminopentylamino,
     dimethylaminopentylamino, ethylaminoethylamino,
     diethylaminoethylamino, ethylaminopropylamino,
80
     diethylaminopropylamino, ethylaminobutylamino,
     diethylaminobutylamino, ethylaminopentylamino,
     methylaminocarbonyl, methylcarbonyl, and ethylcarbonyl;
     or
          a pharmaceutically-acceptable salt or tautomer
85
     thereof.
                A compound of Claim 105 wherein:
          R1 is selected from hydrido, methyl, ethyl,
     hydroxyethyl and propargyl; and
          R2 is R200-piperazinvl-R201 wherein:
 5
          R200 is selected from:
          - (CR202R203) ..-;
          -NR<sup>202</sup>-;
          -S-;
          -0-:
10
          or R200 represents a bond;
          R^{201} represents one or more radicals selected from
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, (1-hydroxy-1,1-
     dimethyl) ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
    cyclohexyl, methoxymethylene, methoxyethylene,
15
     ethoxyethylene, methoxyphenylene, ethoxyphenylene,
    cyclopropylcarbonyl, cyclobutylcarbonyl,
    cyclopentylcarbonyl, cyclohexylcarbonyl, benzoyl,
    chlorobenzoyl, fluorobenzoyl, hydroxymethylcarbonyl,
20
    hydroxyethylcarbonyl, hydroxypropylcarbonyl,
    carboxymethylcarbonyl, carboxyethylcarbonyl,
    carboxypropylcarbonyl, methoxymethylcarbonyl,
    methoxyethylcarbonyl, methoxypropylcarbonyl,
    ethoxymethylcarbonyl, ethoxyethylcarbonyl.
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25
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
     methylsulfonylmethylene, amino, aminomethyl, aminoethyl,
30
     aminopropyl, phenylamino, benzylamino,
     methylaminomethylene, ethylaminomethylene,
     methylaminoethylene, ethylaminoethylene, aminocarbonyl,
     methylcarbonylamino, ethylcarbonylamino,
35
     methylaminomethylcarbonyl, ethylaminomethylcarbonyl,
     methylcarbonylaminomethylene,
     ethylcarbonylaminomethylene,
     aminomethylcarbonylaminocarbonylmethylene,
     methoxycarbonylamino, ethoxycarbonylamino,
40
     methoxymethylcarbonylamino, methoxyethylcarbonylamino,
     ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,
     methoxycarbonylaminomethylene,
     ethoxycarbonylaminomethylene, and methylsulfonylamino;
     and
          \mathbb{R}^{202} and \mathbb{R}^{203} are independently selected from hydrido,
45
     methyl, ethyl, phenyl and benzyl; and
          y is 0, 1 or 2; and
          R4 is phenyl, wherein said phenyl is optionally
     substituted with one or more radicals independently
50
     selected from fluoro, chloro, methyl, ethyl, methoxy and
     ethoxy; and
          R<sup>5</sup> is selected from hydrido, fluoro, chloro, bromo,
     hydroxy, methyl, ethyl, cyano, carboxy, methoxy,
     methoxycarbonyl, aminocarbonyl, acetyl, methylamino,
55
    dimethylamino, ethylamino, dimethylaminoethylamino,
     hydroxyethylamino, hydroxypropylamino, hydroxybutylamino,
    hydroxycyclopropylamino, hydroxycyclobutylamino,
    hydroxycyclopentylamino, hydroxycyclohexylamino, (1-
    ethyl-2-hydroxy) ethylamino, aminomethyl,
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cyclopropylamino, amino, ethoxycarbonylamino,

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methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, methylaminopropylamino, dimethylaminopropylamino, 65 methylaminobutylamino, dimethylaminobutylamino, methylaminopentylamino, dimethylaminopentylamino, ethylaminoethylamino, diethylaminoethylamino, ethylaminopropylamino, diethylaminopropylamino, ethylaminobutylamino, diethylaminobutylamino, 70 ethylaminopentylamino, methylaminocarbonyl, methylcarbonyl, and ethylcarbonyl; or

a pharmaceutically-acceptable salt or tautomer thereof,

107. A compound of Claim 94 having the Formula XA:

wherein:

10

Z represents a carbon atom or a nitrogen atom; and 5 R^1 is selected from hydrido, methyl, ethyl, hydroxyethyl and propargyl; and R^2 is R^{200} -cyclohexyl- R^{201} wherein: R200 is selected from: - (CR202R203),-; -NR²⁰²-; -S-;

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-0-:

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or R200 represents a bond;
          R<sup>201</sup> represents one or more radicals selected from
15
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-
     1,1-dimethyl)ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
     cyclohexyl, methoxymethylene, methoxyethylene,
     methoxypropylene, ethoxyethylene, ethoxypropylene,
20
     propoxyethylene, propoxypropylene, methoxyphenylene,
     ethoxyphenylene, propoxyphenylene, cyclopropylcarbonyl,
     cyclobutylcarbonyl, cyclopentylcarbonyl,
     cyclohexylcarbonyl, benzoyl, chlorobenzovl,
     fluorobenzoyl, hydroxymethylcarbonyl,
25
     hydroxyethylcarbonyl, hydroxypropylcarbonyl,
     carboxymethylcarbonyl, carboxyethylcarbonyl,
     carboxypropylcarbonyl, methoxymethylcarbonyl,
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl,
30
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
35
     methylsulfonylmethylene, amino, aminomethyl, aminoethyl,
     aminopropyl, phenylamino, benzylamino,
     methylaminomethylene, ethylaminomethylene,
    methylaminoethylene, ethylaminoethylene, aminocarbonyl,
    methylcarbonylamino, ethylcarbonylamino,
40
    methylaminomethylcarbonyl, ethylaminomethylcarbonyl,
    methylcarbonylaminomethylene.
     ethylcarbonylaminomethylene,
     aminomethylcarbonylaminocarbonylmethylene,
    methoxycarbonylamino, ethoxycarbonylamino,
45
    methoxymethylcarbonylamino, methoxyethylcarbonylamino,
    ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,
    methoxycarbonylaminomethylene,
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ethoxycarbonylaminomethylene, methylimidocarbonyl, ethylimidocarbonyl, amidino, methylamidino, methylamidino, benzylamidino, guanidino, guanidinomethylene, guanidinoethylene, and methylsulfonylamino; and

R²⁰² and R²⁰³ are independently selected from hydrido, methyl, ethyl, propyl, butyl, phenyl and benzyl; and

y is 0, 1 or 2; and

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R⁴ is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, ethyl, methoxy and ethoxy; and

R⁵ is selected from hydrido, fluoro, chloro, bromo, hydroxy, methyl, ethyl, propyl, benzyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, dimethylamino, 2-methylbutylamino, ethylamino, dimethylaminoethylamino, hydroxyethylamino, hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino, hydroxycyclopentylamino, hydroxycyclobexylamino, imidazolylamino, morpholinylethylamino, (1-ethyl-2-hydroxy)ethylamino, piperidinylamino, pyridinylmethylamino, phenylmethylpiperidinylamino,

pyridinylmethylamino, phenylmethylpiperidinylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, methylaminopropylamino.

dimethylaminoethylamino, methylaminopropylamino, dimethylaminopropylamino, methylaminobutylamino, dimethylaminobutylamino, methylaminopentylamino, dimethylaminopentylamino, ethylaminoethylamino, diethylaminoethylamino, ethylaminopropylamino,

diethylaminopropylamino, ethylaminobutylamino, diethylaminobutylamino, ethylaminopentylamino, methylaminocarbonyl, methylcarbonyl, and ethylcarbonyl; or

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a pharmaceutically-acceptable salt or tautomer
85
     thereof.
           108. A compound of Claim 107 wherein:
           R1 is selected from hydrido, methyl, ethyl,
     hydroxyethyl and propargyl; and
           R<sup>2</sup> is R<sup>200</sup>-cyclohexyl-R<sup>201</sup> wherein:
 5
           R<sup>200</sup> is selected from:
           - (CR202R203),-;
           -NR202-;
           -S-:
           -0-:
10
           or R<sup>200</sup> represents a bond;
          \mathbb{R}^{201} represents one or more radicals selected from
     the group consisting of hydroxy, hydroxymethyl,
     hydroxyethyl, hydroxypropyl, (1-hydroxy-1,1-
     dimethyl)ethyl, cyclopropyl, cyclobutyl, cyclopentyl,
     cyclohexyl, methoxymethylene, methoxyethylene,
15
     methoxypropylene, ethoxyethylene, ethoxypropylene,
     propoxyethylene, propoxypropylene, methoxyphenylene,
     ethoxyphenylene, propoxyphenylene, cyclopropylcarbonyl,
     cyclobutylcarbonyl, cyclopentylcarbonyl,
     cyclohexylcarbonyl, benzoyl, chlorobenzoyl,
20
     fluorobenzoyl, hydroxymethylcarbonyl,
     hydroxyethylcarbonyl, hydroxypropylcarbonyl,
     carboxymethylcarbonyl, carboxyethylcarbonyl,
     carboxypropylcarbonyl, methoxymethylcarbonyl,
25
     methoxyethylcarbonyl, methoxypropylcarbonyl,
     ethoxymethylcarbonyl, ethoxyethylcarbonyl,
     ethoxypropylcarbonyl, propoxymethylcarbonyl,
     propoxyethylcarbonyl, propoxypropylcarbonyl,
     methoxyphenylcarbonyl, ethoxyphenylcarbonyl,
30
     propoxyphenylcarbonyl, piperidinylmethylcarbonyl,
     piperazinylmethylcarbonyl, morpholinylcarbonyl,
     methylsulfonylmethylene, amino, aminomethyl, aminoethyl,
     aminopropyl, phenylamino, benzylamino,
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methylaminomethylene, ethylaminomethylene,
methylaminoethylene, ethylaminoethylene, aminocarbonyl,
methylcarbonylamino, ethylcarbonylamino,
methylaminomethylcarbonyl, ethylaminomethylcarbonyl,
methylcarbonylaminomethylene,
ethylcarbonylaminomethylene,
aminomethylcarbonylaminocarbonylmethylene,
methoxycarbonylamino, ethoxycarbonylamino,
methoxymethylcarbonylamino, methoxyethylcarbonylamino,
ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,

45 ethoxycarbonylaminomethylene; and

methoxycarbonylaminomethylene, and

 R^{202} and R^{203} are independently selected from hydrido, methyl, ethyl, phenyl and benzyl; and

y is 0, 1 or 2; and

50

R⁴ is phenyl, wherein said phenyl is optionally substituted with one or more radicals independently selected from fluoro, chloro, methyl, ethyl, methoxy and ethoxy; and

R5 is selected from hydrido, fluoro, chloro, bromo, hydroxy, methyl, ethyl, cyano, carboxy, methoxy, methoxycarbonyl, aminocarbonyl, acetyl, methylamino, 55 dimethylamino, ethylamino, dimethylaminoethylamino, hydroxyethylamino, hydroxypropylamino, hydroxybutylamino, hydroxycyclopropylamino, hydroxycyclobutylamino, hydroxycyclopentylamino, hydroxycyclohexylamino, (1-60 ethyl-2-hydroxy) ethylamino, aminomethyl, cyclopropylamino, amino, ethoxycarbonylamino, methoxyphenylmethylamino, phenylmethylamino, fluorophenylmethylamino, fluorophenylethylamino, methylaminoethylamino, dimethylaminoethylamino, 65 methylaminopropylamino, dimethylaminopropylamino, methylaminobutylamino, dimethylaminobutylamino, methylaminopentylamino, dimethylaminopentylamino, ethylaminoethylamino, diethylaminoethylamino,

ethylaminopropylamino, diethylaminopropylamino,

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70
     ethylaminobutylamino, diethylaminobutylamino.
     ethylaminopentylamino, methylaminocarbonyl,
     methylcarbonyl, and ethylcarbonyl; or
           a pharmaceutically-acceptable salt or tautomer
     thereof.
           109. A compound of Claim 107 wherein:
           R1 is hydrido; and
           R^2 is R^{200}-cyclohexyl-R^{201} wherein:
           R<sup>200</sup> is selected from:
 5
           methylene;
           -NR<sup>202</sup>-;
           -S-;
           -0-;
           or R200 represents a bond;
           R^{201} represents one or more radicals selected from
10
     the group consisting of amino, aminomethyl, aminoethyl,
     aminopropyl, phenylamino, benzylamino,
     methylaminomethylene, ethylaminomethylene,
     methylaminoethylene, ethylaminoethylene, aminocarbonyl,
15
     methylcarbonylamino, ethylcarbonylamino,
     methylaminomethylcarbonyl, ethylaminomethylcarbonyl,
     methylcarbonylaminomethylene,
     ethylcarbonylaminomethylene,
     aminomethylcarbonylaminocarbonylmethylene,
     methoxycarbonylamino, ethoxycarbonylamino,
20
     methoxymethylcarbonylamino, methoxyethylcarbonylamino,
     ethoxymethylcarbonylamino, ethoxyethylcarbonylamino,
     methoxycarbonylaminomethylene, and
     ethoxycarbonylaminomethylene; and
          R^{202} is selected from hydrido, methyl, phenyl and
25
     benzyl; and
          R4 is phenyl, wherein said phenyl is optionally
     substituted with one or more radicals independently
     selected from fluoro, chloro, methyl, and methoxy; and
30
          R<sup>5</sup> is selected from hydrido, methylamino,
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dimethylamino, 2-methylbutylamino, ethylamino, dimethylaminoethylamino, hydroxypropylamino, hydroxyethylamino, hydroxypropylamino, hydroxyethylamino, hydroxyeyclobutylamino, hydroxyeyclopropylamino, hydroxyeyclobutylamino, (1-ethyl-2-hydroxy)ethylamino, aminomethyl, cyclopropylamino, amino, dimethylaminoethylamino, dimethylaminopropylamino, dimethylaminobutylamino, dimethylaminopropylamino, diethylaminoethylamino, diethylaminopropylamino, diethylaminobutylamino, and diethylaminopentylamino; or

a pharmaceutically-acceptable salt or tautomer thereof.

110. A compound of Claim 94 wherein R^2 comprises a substituted piperidinyl or piperazinyl moiety with at least one substituent attached to the distal nitrogen heteroatom or to a carbon ring atom adjacent to the distal nitrogen heteroatom of the piperidine or piperazine ring.

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5

- 111. A compound Claim 94 wherein R² comprises a substituted piperidinyl moiety with at least one substituent attached to the distal nitrogen heteroatom or to a carbon ring atom adjacent to the distal nitrogen heteroatom of the piperidine ring.
- 112. A compound of Claim 94 wherein R^2 comprises a substituted piperazinyl moiety with at least one substituent attached to the distal nitrogen heteroatom or to a carbon ring atom adjacent to the distal nitrogen heteroatom of the piperazine ring.
- 113. A compound of Claim 94 wherein Z represents a carbon atom.

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- 114. A compound of Claim 94 wherein Z represents a nitrogen atom.
 - 115. A compound of Claim 94 wherein R1 is hydrido.
- 116. A compound of Claim 94 wherein R^{200} represents a bond.
- 117. A compound of Claim 94 wherein R²⁰¹ represents one or more radicals selected from the group consisting of lower hydroxyalkyl, lower hydroxyalkylcarbonyl, and lower alkylaminoalkylene.
- 118. A compound of Claim 94 wherein R²⁰¹ represents one or more radicals selected from the group consisting of hydroxymethyl, hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-1,1-dimethyl)ethyl, hydroxymethylcarbonyl, hydroxyethylcarbonyl, hydroxypropylcarbonyl, methylaminomethylene, ethylaminomethylene, methylaminoethylene, and ethylaminoethylene.

- 119. A compound of Claim 94 wherein \mathbb{R}^4 is optionally substituted phenyl.
- 120. A compound of Claim 94 wherein R⁴ is phenyl optionally substituted at a substitutable position with one or more radicals independently selected from chloro, fluoro, bromo and iodo.
- 121. A compound of Claim 94 wherein \mathbb{R}^4 is phenyl optionally substituted at the meta or para position with one or more chloro radicals.
 - 122. A compound of Claim 94 wherein R⁵ is hydrido.

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123. A compound of Claim 94 wherein:

R1 is hydrido;

R²⁰⁰ represents a bond;

 $$\rm R^{201}$$ represents one or more radicals selected from the group consisting of lower hydroxyalkyl, lower

hydroxyalkylcarbonyl, and lower alkylaminoalkylene.

R' is phenyl optionally substituted at a substitutable position with one or more radicals independently selected from halo; and

10 R⁵ is hydrido.

5

10

124. A compound of Claim 94 wherein:

R1 is hydrido;

R²⁰⁰ represents a bond;

R²⁰¹ represents one or more radicals selected from the group consisting of hydroxymethyl, hydroxyethyl, hydroxypropyl, hydroxybutyl, (1-hydroxy-1,1-

nydroxypropy1, nydroxybuty1, (1-nydroxy-1,1-dimethyl)ethyl, hydroxymethylcarbonyl, hydroxypropylcarbonyl,

methylaminomethylene, ethylaminomethylene, methylaminoethylene, and ethylaminoethylene;

 $\rm R^4$ is phenyl optionally substituted at a substitutable position with one or more radicals independently selected from chloro, fluoro, bromo and iodo; and

15 R⁵ is hydrido.

125. A compound selected from compounds, their tautomers and their pharmaceutically acceptable salts, of the group consisting of:

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1160

126. A compound of Formula IA

wherein

5

10

R¹ is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkoxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl,

alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl,

alkylthioalkylene, alkenylthioalkylene,

20 heterocyclylsulfonyl, alkylaminoalkylene,
 alkylsulfonylalkylene, acyl, acyloxycarbonyl,
 alkoxycarbonylalkylene, aryloxycarbonylalkylene,
 heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene,
 aryloxycarbonylarylene, heterocyclyloxycarbonylarylene,
25 alkylcarbonylalkylene, arylcarbonylalkylene,

alkylcarbonylalkylene, arylcarbonylalkylene, heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,

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30 arylcarbonyloxyarylene, and heterocyclylcarbonyloxyarylene; or

R1 has the formula

wherein:

40

i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl,

aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclyl, alkylheterocyclylalkylene, alkylheterocyclylarylene,

aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene,

aryloxycarbonylarylene, alkylaryloxycarbonylarylene, arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene,

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alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, 65 cycloalkylthioalkylene, alkylthioarylene, aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, 70 alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene. arylthioalklylarylene, and alkylsulfonylarylene groups may be optionally substituted with one or more radicals 75 independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or

R²⁷ is -CHR²⁸R²⁹ wherein R²⁸ is alkoxycarbonyl, and R²⁹ is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl and nitro; or

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R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy; and

```
R2 is R200-cvcloalkyl-R201 wherein:
100
              R<sup>200</sup> is selected from:
              - (CR202R203),-;
              -C(O)-;
              -C(O)-(CH2),-;
              -C(O)-O-(CH2),-;
105
              -(CH_2)_v-C(O)-;
              -O-(CH<sub>2</sub>),-C(O)-;
              -NR<sup>202</sup>-;
              -NR^{202} - (CH_2)_{v} - ;
              - (CH<sub>2</sub>),-NR<sup>202</sup>-;
              -(CH_2)_v-NR^{202}-(CH_2)_z-;
110
              -(CH_2)_V - C(O) - NR^{202} - (CH_2)_V - i
              -(CH_2)_v - NR^{202} - C(O) - (CH_2)_z - ;
             -(CH_2)_v - NR^{202} - C(O) - NR^{203} - (CH_2)_z - ;
             -S(O) x- (CR202R203) y-;
115
             -(CR^{202}R^{203})_{v}-S(O)_{x}-;
             -S(O) - (CR202R203) -O-;
             -S(O)_{x}-(CR^{202}R^{203})_{y}-C(O)-;
             -O- (CH2) v-;
             - (CH<sub>2</sub>) -O-;
120
             -S-; and
             -0-:
             \mathbb{R}^{201} represents one or more radicals selected from
       the group consisting of hydrido, halogen, hydroxy,
       carboxy, keto, alkyl, hydroxyalkyl, haloalkyl,
       cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,
125
       aralkyl, heterocyclylalkylene, alkylcarbonyl,
       hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl,
       haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene,
       alkoxycarbonyl, carboxyalkylcarbonyl,
130
       alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,
       alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl,
       alkylamino, aralkylamino, alkylaminoalkylene,
       aminocarbonyl, alkylcarbonylamino,
       alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl,
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alkylaminoalkylcarbonylamino,
aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino,
alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene,
alkylimidocarbonyl, amidino, alkylamidino,
aralkylamidino, guanidino, guanidinoalkylene, and
alkylsulfonylamino; and

 \mathbb{R}^{202} and \mathbb{R}^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

y and z are independently 0, 1, 2, 3, 4, 5 or 6 wherein y + z is less than or equal to 6; and

x is 0, 1 or 2; and

R³ is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylakyl, thiazolylamino,

150

145

wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylakyl, thiazolylamino,

155

160

groups may be optionally substituted with one or more radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkenylamino, alkynylamino, cycloalkylamino, cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy,

1165

165 hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino, 170 alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino. heterocyclylheterocyclylalkylamino, 175 alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR44R45 wherein R44 is alkylcarbonyl or amino, and R45 is alkyl or 180 aralkyl; and R4 is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R^4 is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, 185 alkylthioalkylene, arylthioalkylene, alkylsulfinyl, alkylsulfinylalkylene, arylsulfinylalkylene, alkylsulfonyl, alkylsulfonylalkylene, arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, 190 alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano, nitro, alkylamino, arylamino, alkylaminoalkylene, arylaminoalkylene, aminoalkylamino, and hydroxy; provided R3 is not 2-pyridinyl when R4 is a phenyl

provided R' is not 2-pyridinyl when R' is a phenyl 195 ring containing a 2-hydroxy substituent and when R' is hydrido; and

further provided that R^4 is not methylsulfonylphenyl or aminosulfonylphenyl; and

further provided that R^1 is not methylsulfonylphenyl;

200 or

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a pharmaceutically-acceptable salt or tautomer thereof.

127. A compound of Formula IA

wherein

5 R¹ is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene,

15 alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkenylamin

alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoslkylene

20 heterocyclylsulfonyl, alkylaminoalkylene, alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, aryloxycarbonylalkylene,

25 alkylcarbonylalkylene, arylcarbonylalkylene, heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene,

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alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene, arylcarbonyloxyarylene, and heterocyclylcarbonyloxyarylene; or

R1 has the formula

wherein:

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i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclylalkylene, alkylheterocyclylarylene, aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene,

alkoxyarylene, alkoxyarylene, alkoxyarylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene,

60 aryloxycarbonylarylene, alkylaryloxycarbonylarylene,

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arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, 65 cycloalkylthioalkylene, alkylthioarylene, aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, 70 alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene. aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups 75 may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or R^{27} is $-CHR^{28}R^{29}$ wherein R^{28} is alkoxycarbonyl, and R^{29} is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, 80 alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl 85 and nitro; or

R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be

90

95 heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals

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independently selected from halogen, alkyl and alkoxy;
        and
              R^2 is R^{200}-aryl-R^{201} wherein:
              R<sup>200</sup> is selected from:
100
              - (CR202R203),-;
              -C(0) -;
              -C(O)-(CH<sub>2</sub>),-;
              -C(O)-O-(CH2),-;
105
              - (CH<sub>2</sub>),-C(O)-;
              -O- (CH<sub>2</sub>)<sub>v</sub>-C(O) -;
              -NR202-:
              -NR^{202}-(CH_2)_{v}-;
              - (CH<sub>2</sub>) ,-NR<sup>300</sup>-;
              -(CH_2)_v-NR^{202}-(CH_2)_{z1}-;
110
              -(CH_2)_v - C(O) - NR^{202} - (CH_2)_z - i
              -(CH_2)_v-NR^{202}-C(O)-(CH_2)_z-;
              -(CH_2)_v-NR^{202}-C(O)-NR^{203}-(CH_2)_z-;
              -S(O),-(CR202R203),-;
115
              - (CR202R203),-S(O),-;
              -S (O) x- (CR202R203) v-O-;
              -S(O)_{x}-(CR^{202}R^{203})_{y}-C(O)-;
              -O- (CH<sub>2</sub>),-;
              -(CH_2)_v - O - ; and
120
              -0-:
              {\ensuremath{\mathbb{R}}}^{201} represents one or more radicals selected from
       the group consisting of hydrido, halogen, hydroxy,
       carboxy, keto, alkyl, hydroxyalkyl, haloalkyl,
       cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,
       aralkyl, heterocyclylalkylene, alkylcarbonyl,
125
       hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl,
       haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene,
       alkoxycarbonyl, carboxyalkylcarbonyl.
       alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,
       alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl,
130
       alkylamino, aralkylamino, alkylaminoalkylene,
       aminocarbonyl, alkylcarbonylamino,
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1170

alkylcarbonylaminoalkylcarbonyl, alkylaminoalkylcarbonyl, alkylaminoalkylcarbonylamino,

aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino, alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene, alkylimidocarbonyl, amidino, alkylamidino, aralkylamidino, guanidino, guanidinoalkylene, and alkylsulfonylamino; and

 R^{202} and R^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

R³⁰⁰ is selected from alkyl, aryl and aralkyl; and
y and z are independently 0, 1, 2, 3, 4, 5 or 6
wherein y + z; and yl is 1, 2, 3, 4, 5 or 6; wherein y +
z and yl + z are less than or equal to 6; and

x is 0, 1 or 2; and

R³ is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

$$\begin{bmatrix} \\ \\ \\ \\ \\ \end{bmatrix} \qquad ; \qquad \begin{bmatrix} \\ \\ \\ \\ \\ \end{bmatrix} \qquad ; \text{ and } \begin{bmatrix} \\ \\ \\ \\ \\ \end{bmatrix}$$

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wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

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groups may be optionally substituted with one or more radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino,

1171

alkenylamino, alkynylamino, cycloalkylamino, cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, 165 hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, 170 aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino, 175 heterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and $-NR^{44}R^{45}$ 180 wherein R44 is alkylcarbonyl or amino, and R45 is alkyl or aralkyl; and R4 is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R^4 is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, 185 alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl, alkylsulfinylalkylene, arylsulfinylalkylene, alkylsulfonyl, alkylsulfonylalkylene, 190 arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano, nitro, alkylamino, arylamino, alkylaminoalkylene, arylaminoalkylene, aminoalkylamino, and hydroxy; 195 provided R3 is not 2-pyridinyl when R4 is a phenyl

hydrido; and further provided that R4 is not methylsulfonylphenyl

ring containing a 2-hydroxy substituent and when R^1 is

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or aminosulfonylphenyl; and

further provided that R¹ is not methylsulfonylphenyl; or

a pharmaceutically-acceptable salt or tautomer thereof.

128. A compound of Formula IA

wherein

R¹ is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl,

hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene,

alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl,

20 heterocyclylsulfonyl, alkylaminoalkylene, alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene,

1173

alkylcarbonylalkylene, arylcarbonylalkylene,
heterocyclylcarbonylalkylene, alkylcarbonylarylene,
arylcarbonylarylene, heterocyclylcarbonylarylene,
alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene,
heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,
arylcarbonyloxyarylene, and
heterocyclylcarbonyloxyarylene; or

R1 has the formula

wherein:

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i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and R²⁷ is selected from alkyl, cycloalkyl, alkynyl,

aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene,

alkylaralkyl, aralkylarylene, alkylheterocyclyl, alkylheterocyclylalkylene, alkylheterocyclylarylene,

aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene,

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arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, alkylaryloxycarbonylarylene, 60 arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, 65 cycloalkylthioalkylene, alkylthioarylene, aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, 70 alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups 75 may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or R^{27} is $-CHR^{28}R^{29}$ wherein R^{28} is alkoxycarbonyl, and R^{29} is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, 80 alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl 85 and nitro: or

R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and

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alkoxycarbonylamino; wherein said aryl,
 95
       heterocyclylalkylene and aryloxyalkylene radicals may be
       optionally substituted with one or more radicals
       independently selected from halogen, alkyl and alkoxy;
       and
             R^2 is R^{200}-heterocyclyl-R^{201} wherein:
100
             R<sup>200</sup> is selected from:
             - (CR301R302),-;
             -C(O)-(CH<sub>2</sub>)<sub>1/2</sub>-;
             -C(O)-O-(CH2),-;
             - (CH<sub>2</sub>),-C(O)-;
105
             -O- (CH<sub>2</sub>),-C(O)-;
             -NR^{303}-;
             -NR^{303} - (CH_2)_{v} - ;
             -(CH_2)_{11}-NR^{202}-;
             -(CH_2)_v-NR^{202}-(CH_2)_{z1}-;
110
             -(CH_2)_v - C(O) - NR^{202} - (CH_2)_v - i
             -(CH_2)_v - NR^{202} - C(O) - (CH_2)_z - ;
             -(CH_2)_{v}-NR^{202}-C(O)-NR^{203}-(CH_2)_{z}-;
             -S(O) - (CR202R203) -;
             - (CR202R203),-S(O),-;
115
             -S(0),-(CR202R203),-O-;
             -S(O)_{x}-(CR^{202}R^{203})_{y}-C(O)-;
             -0-(CH_2)_v-; and
             - (CH<sub>2</sub>),-O-;
             R^{201} represents one or more radicals selected from
120
       the group consisting of hydrido, halogen, hydroxy,
       carboxy, keto, alkyl, hydroxyalkyl, haloalkyl,
       cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,
       aralkyl, heterocyclylalkylene, alkylcarbonyl,
       hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl,
      haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene,
125
       alkoxycarbonyl, carboxyalkylcarbonyl,
       alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,
       alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl,
       alkylamino, aralkylamino, alkylaminoalkylene,
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aminocarbonyl, alkylcarbonylamino,
 alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl,
 alkylaminoalkylcarbonylamino,
 aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino,
 alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene,
 alkylimidocarbonyl, amidino, alkylamidino,
 aralkylamidino, guanidino, guanidinoalkylene, and
 alkylsulfonylamino; and

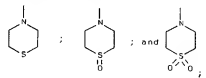
 R^{202} and R^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

140 R³⁰¹ and R³⁰² are independently selected from aryl and aralkyl; and

R³⁰³ is selected from alkyl, aryl and aralkyl; and y and z are independently 0, 1, 2, 3, 4, 5 or 6; and yl is 1, 2, 3, 4, 5 or 6; wherein y + z and yl + z are less than or equal to 6; and

x is 0, 1 or 2; wherein either x or y is other than 0 when R^{200} is $-S(O)_x-(CR^{202}R^{203})_y-$; and

R³ is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,



wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl,

155 thiazolylalkyl, thiazolylamino,

145

150

groups may be optionally substituted with one or more radicals independently selected from halo, keto, alkyl,

1177

160 aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkenylamino, alkynylamino, cycloalkylamino, 165 cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, 170 aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino, 175 heterocyclylheterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arvlhydrazinyl, and -NR44R45 180 wherein R44 is alkylcarbonyl or amino, and R45 is alkyl or aralkyl; and R4 is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein $\ensuremath{\mathrm{R}}^4$ is optionally substituted with one or more radicals 185 independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl, alkylsulfinylalkylene, arylsulfinylalkylene, 190 alkylsulfonyl, alkylsulfonylalkylene, arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano, nitro, alkylamino, arylamino, alkylaminoalkylene,

arylaminoalkylene, aminoalkylamino, and hydroxy;

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provided R^3 is not 2-pyridinyl when R^4 is a phenyl ring containing a 2-hydroxy substituent and when R^1 is hydrido; and

further provided R^2 is selected from aryl, heterocyclyl, unsubstituted cycloalkyl and cycloalkenyl when R^4 is hydrido; and

further provided that R4 is not methylsulfonylphenyl or aminosulfonylphenyl; and

further provided that \mathbb{R}^1 is not methylsulfonylphenyl; 205 or

a pharmaceutically-acceptable salt or tautomer thereof.

129. A compound of Formula IA

wherein

200

R¹ is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkylene, alkylthioalkylene, alkylthioalkylene, alkylthioalkylene, alkynylamino, arylamino, heterocyclylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino,

alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl,

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arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, 20 heterocyclylsulfonyl, alkylaminoalkylene, alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, arylcarbonylalkylene, 25 heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene, 30 arylcarbonyloxyarylene, and heterocyclylcarbonyloxyarylene; or

R1 has the formula

wherein:

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i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclylalkylene, alkylheterocyclylarylene, aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene,

1180

alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene. alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl, alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, 55 alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene, 60 aryloxycarbonylarylene, alkylaryloxycarbonylarylene, arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, 65 cycloalkylthioalkylene, alkylthioarylene, aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, 70 heterocyclylalkylene, alkylheterocyclylarylene, alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups 75 may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or $\mbox{R}^{27} \mbox{ is -CHR}^{28}\mbox{R}^{29} \mbox{ wherein } \mbox{R}^{28} \mbox{ is alkoxycarbonyl, and } \mbox{R}^{29}$ is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, 80 alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl 85 and nitro; or

 R^{26} and R^{27} together with the nitrogen atom to which

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they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl,

- heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl,
- 95 heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy; and
- R² is selected from hydrido, halogen, mercapto,
 alkyl, alkenyl, alkynyl, aryl, heterocyclyl, haloalkyl,
 hydroxyalkyl, aralkyl, alkylheterocyclyl,
 heterocyclylalkyl, heterocyclylheterocyclyl,
 heterocyclylalkylheterocyclyl, alkylamino, alkenylamino,
 alkynylamino, arylamino, aryl(hydroxyalkyl)amino,
- heterocyclylamino, heterocyclylalkylamino, aralkylamino, N-alkyl-N-alkynyl-amino, aminoalkyl, aminoaryl, aminoalkylamino, aminocarbonylalkylene, arylaminoalkylene, alkylaminoalkylene, arylaminoarylene, alkylaminoarylene, alkylaminoalkylamino,
- alkylcarbonylaminoalkylene,
 aminoalkylcarbonylaminoalkylene,
 alkylaminoalkylcarbonylamino, cycloalkyl, cycloalkenyl,
 aminoalkylthio, alkylaminocarbonylalkylthio,
 alkylaminoalkylaminocarbonylalkylthio, alkoxy,
- heterocyclyloxy, alkylthio, cyanoalkylthio, alkenylthio, alkynylthio, carboxyalkylthio, arylthio, heterocyclylthio, alkoxycarbonylalkylthio, alkylsulfinyl, alkylsulfonyl, carboxy, carboxyalkyl, alkoxyalkyl, alkoxyalkylthio, carboxycycloalkyl, carboxycycloalkenyl,
- carboxyalkylamino, alkoxycarbonyl, heterocyclylcarbonyl, alkoxycarbonylalkyl, alkoxycarbonylalkylamino, alkoxycarbonylheterocyclyl,

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alkoxycarbonylheterocyclylcarbonyl, alkoxyalkylamino,
       alkoxycarbonylaminoalkylene, alkoxycarbonylaminoalkoxy,
       alkoxycarbonylaminoalkylamino, heterocyclylsulfonyl,
125
       aralkythio, heterocyclylalkylthio, aminoalkoxy,
        cyanoalkoxy, carboxyalkoxy, aryloxy, aralkoxy,
       alkenyloxy, alkynyloxy, and heterocyclylalkyloxy; wherein
       the aryl, heterocyclyl, heterocyclylalkyl, cycloalkyl and
       cycloalkenyl groups may be optionally substituted with
130
       one or more radicals independently selected from halo,
       keto, amino, alkyl, alkenyl, alkynyl, aryl, heterocyclyl,
       aralkyl, heterocyclylalkyl, epoxyalkyl,
       amino(hydroxyalkyl) carboxy, alkoxy, aryloxy, aralkoxy,
       haloalkyl, alkylamino, alkynylamino,
135
       alkylaminoalkylamino, heterocyclylalkylamino,
       alkylcarbonyl, alkoxycarbonyl, alkylsulfonyl,
       arylsulfonyl, and aralkylsulfonyl; or
              R^2 is R^{200}-heterocyclyl-R^{201}, R^{200}-aryl-R^{201}, or R^{200}-
140
       cycloalkyl-R201 wherein:
             R<sup>200</sup> is selected from:
              - (CR202R203),-;
              -C(0)-;
              -C(O)-(CH2)v-;
145
             -C(O)-O-(CH<sub>2</sub>)<sub>v</sub>-;
              - (CH<sub>2</sub>),-C(O)-;
             -O-(CH<sub>2</sub>),-C(O)-;
             -NR<sup>202</sup>-;
             -NR^{202} - (CH_2)_{v} - ;
150
             - (CH<sub>2</sub>),-NR<sup>202</sup>-;
             -(CH_2)_v - NR^{202} - (CH_2)_z - ;
             - (CH<sub>2</sub>),-C(O)-NR<sup>202</sup>-(CH<sub>2</sub>),-:
             -(CH_2)_v-NR^{202}-C(O)-(CH_2)_z-;
             -(CH_2)_v - NR^{202} - C(O) - NR^{203} - (CH_2)_v - :
155
             -S(0)_{x}-(CR^{202}R^{203})_{y}-;
             -(CR^{202}R^{203})_y-S(O)_{x^-};
             -S(O),-(CR202R203),-O-;
             -S(O)_{x}-(CR^{202}R^{203})_{y}-C(O)-;
```

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-O- (CH<sub>2</sub>),-;
160
            - (CH<sub>2</sub>),-O-;
            -S-;
            -0-;
            or R200 represents a bond:
            R^{201} represents one or more radicals selected from
165
       the group consisting of hydrido, halogen, hydroxy,
       carboxy, keto, alkyl, hydroxyalkyl, haloalkyl,
       cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,
       aralkyl, heterocyclylalkylene, alkylcarbonyl,
       hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl,
170
       haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene,
       alkoxycarbonyl, carboxyalkylcarbonyl,
       alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,
       alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl,
      alkylamino, aralkylamino, alkylaminoalkylene,
      aminocarbonyl, alkylcarbonylamino,
175
       alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl.
      alkylaminoalkylcarbonylamino,
      aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino,
      alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene,
180
      alkylimidocarbonyl, amidino, alkylamidino,
      aralkylamidino, guanidino, guanidinoalkylene, and
      alkylsulfonylamino; and
            R^{202} and R^{203} are independently selected from hydrido,
      alkyl, aryl and aralkyl; and
            y and z are independently 0, 1, 2, 3, 4, 5 or 6
185
      wherein y + z is less than or equal to 6; and
            x is 0, 1 or 2; or
            R<sup>2</sup> is -NHCR<sup>204</sup>R<sup>205</sup> wherein R<sup>204</sup> is alkylaminoalkylene,
      and R<sup>205</sup> is aryl; or
            \mbox{R}^2 is -\mbox{C(NR}^{206})\,\mbox{R}^{207} wherein \mbox{R}^{206} is selected from
190
      hydrogen and hydroxy, and R<sup>207</sup> is selected from alkyl,
      aryl and aralkyl; or
            R2 has the formula:
```

195 wherein:

j is an integer from 0 to 8; and m is 0 or 1; and

R³⁰ and R³¹ are independently selected from hydrogen, alkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene,

aminoalkyl, alkylaminoalkyl, aminocarbonylalkyl, alkoxyalkyl, and alkylcarbonyloxyalkyl; and

R³² is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene;

 R^{33} is selected from hydrogen, alkyl, $-C(0)\,R^{35},$ $-C(0)\,OR^{35},$ $-SO_2R^{36},$ $-C(0)\,NR^{37}R^{38},$ and $-SO_2NR^{39}R^{40},$ wherein $R^{35},$ $R^{36},$ $R^{37},$ $R^{38},$ R^{39} and R^{40} are independently selected from hydrocarbon, heterosubstituted hydrocarbon and heterocyclyl; and

R³⁴ is selected from hydrogen, alkyl, aminocarbonyl, alkylaminocarbonyl, and arylaminocarbonyl; or

 \mbox{R}^2 is $-\mbox{CR}^{41}\mbox{R}^{42}$ wherein \mbox{R}^{41} is aryl, and \mbox{R}^{42} is hydroxy;

215 and

205

210

 \mathbb{R}^3 is selected from maleimidyl, pyridonyl, thiazolyl, thiazolylakyl, thiazolylamino,

wherein the R³ maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

groups may be optionally substituted with one or more 225 radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkenylamino, alkynylamino, cycloalkylamino, 230 cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, 235 alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, 240 alkylheterocyclylalkylamino, aralkylheterocyclylamino, heterocyclylheterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, 245 haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl,

hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR44R45
wherein R44 is alkylcarbonyl or amino, and R45 is alkyl or
aralkyl; and
R4 is selected from hydrido, alkyl, alkenyl, alkynyl,
cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein

cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R4 is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl,

1186

alkylsulfinylalkylene, arylsulfinylalkylene,
alkylsulfonyl, alkylsulfonylalkylene,
arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy,
aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl,
alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano,
nitro, alkylamino, arylamino, alkylaminoalkylene,
arylaminoalkylene, aminoalkylamino, and hydroxy;
provided R³ is not

 $\bigcup_{N=1}^{N} O \qquad \qquad \bigcup_{N=0}^{N} O$

(IV) (V)

wherein R⁴³ is selected from hydrogen, alkyl, aminoalkyl, alkoxyalkyl, alkenoxyalkyl, and aryloxyalkyl; and

further provided R^2 is selected from aryl, heterocyclyl, unsubstituted cycloalkyl and cycloalkenyl when R^4 is hydrido; and

further provided that R^4 is not methylsulfonylphenyl or aminosulfonylphenyl; and

further provided that R^1 is not methylsulfonylphenyl; or

a pharmaceutically-acceptable salt or tautomer thereof.

130. A compound of Formula IA

1187

wherein

5 R1 is selected from hydrido, hydroxy, alkyl, cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, 10 arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino, 15 alkenylamino, alkynylamino, arylamino, heterocyclylamino, alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene, 20 alkylsulfonylalkylene, acyl, acyloxycarbonyl, alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, 25 alkylcarbonylalkylene, arylcarbonylalkylene, heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,

R1 has the formula

heterocyclylcarbonyloxyarylene; or

arylcarbonyloxyarylene, and

1188

$$\begin{array}{c|c}
 & R^{25} \\
 & \downarrow \\
 & C \\
 & C \\
 & H
\end{array}$$

$$\begin{array}{c|c}
 & Q^{26} \\
 & Q^{27} \\
 & R^{27}
\end{array}$$
(II)

wherein:

40

i is an integer from 0 to 9;

R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclyl, alkylheterocyclylalkylene, alkylheterocyclylarylene,

aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene, aryloxyarylene, aralkoxyarylene, alkoxyheterocyclylalkylene, aryloxyalkoxyarylene, alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl, alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene,

aryloxycarbonylarylene, alkylaryloxycarbonylarylene, arylcarbonylarylene, alkylarylcarbonylarylene, alkoxycarbonylheterocyclylarylene, alkoxycarbonylalkoxylarylene, heterocyclylcarbonylalkylarylene, alkylthioalkylene, cycloalkylthioalkylene, alkylthioarylene,

1189

aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, alkoxy, keto, amino, nitro, and cyano; or

70

75

80

85

R²⁷ is -CHR²⁸R²⁹ wherein R²⁸ is alkoxycarbonyl, and R²⁹ is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, alkoxycarbonylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl and nitro; or

R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl,

90 heterocyclyl, heterocyclylalkylene, alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl,

95 heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy; and

R² is selected from hydrido, halogen, mercapto,
100 alkyl, alkenyl, alkynyl, aryl, heterocyclyl, haloalkyl,
hydroxyalkyl, aralkyl, alkylheterocyclyl,

1190

heterocyclylalkyl, heterocyclylheterocyclyl, heterocyclylalkylheterocyclyl, alkylamino, alkenylamino, alkynylamino, arylamino, aryl(hydroxyalkyl)amino, 105 heterocyclylamino, heterocyclylalkylamino, aralkylamino, N-alkyl-N-alkynyl-amino, aminoalkyl, aminoaryl. aminoalkylamino, aminocarbonylalkylene, arylaminoalkylene, alkylaminoalkylene, arylaminoarylene, alkylaminoarylene, alkylaminoalkylamino, 110 alkylcarbonylaminoalkylene, aminoalkylcarbonylaminoalkylene, alkylaminoalkylcarbonylamino, cycloalkyl, cycloalkenyl, aminoalkylthio, alkylaminocarbonylalkylthio, alkylaminoalkylaminocarbonylalkylthio, alkoxy, heterocyclyloxy, alkylthio, cyanoalkylthio, alkenylthio, 115 alkynylthio, carboxyalkylthio, arylthio, heterocyclylthio, alkoxycarbonylalkylthio, alkylsulfinyl, alkylsulfonyl, carboxy, carboxyalkyl, alkoxyalkyl, alkoxyalkylthio, carboxycycloalkyl, carboxycycloalkenyl, carboxyalkylamino, alkoxycarbonyl, heterocyclylcarbonyl, 120 alkoxycarbonylalkyl, alkoxycarbonylalkylamino, alkoxycarbonylheterocyclyl, alkoxycarbonylheterocyclylcarbonyl, alkoxyalkylamino, alkoxycarbonylaminoalkylene, alkoxycarbonylaminoalkoxy, alkoxycarbonylaminoalkylamino, heterocyclylsulfonyl, 125 aralkythio, heterocyclylalkylthio, aminoalkoxy, cyanoalkoxy, carboxyalkoxy, aryloxy, aralkoxy, alkenyloxy, alkynyloxy, and heterocyclylalkyloxy; wherein the aryl, heterocyclyl, heterocyclylalkyl, cycloalkyl and 130 cycloalkenyl groups may be optionally substituted with one or more radicals independently selected from halo, keto, amino, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, aralkyl, heterocyclylalkyl, epoxyalkyl, amino(hydroxyalkyl) carboxy, alkoxy, aryloxy, aralkoxy, 135 haloalkyl, alkylamino, alkynylamino, alkylaminoalkylamino, heterocyclylalkylamino,

alkylcarbonyl, alkoxycarbonyl, alkylsulfonyl,

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arylsulfonyl, and aralkylsulfonyl; or
              R^2 is R^{200}-heterocyclyl-R^{201}, R^{200}-aryl-R^{201}, or R^{200}-
140
        cycloalkyl-R201 wherein:
              R<sup>200</sup> is selected from:
              - (CR202R203),-;
              -C(0)-;
              -C(O)-(CH2),-;
145
              -C(O)-O-(CH<sub>2</sub>),-;
              - (CH<sub>2</sub>),-C(O)-;
              -O- (CH2) v-C(O) -;
              -NR^{202}-;
              -NR^{202}-(CH_2)_{v}-;
150
              -(CH_2)_v - NR^{202} - ;
              - (CH<sub>2</sub>),-NR<sup>202</sup>-(CH<sub>2</sub>)<sub>z</sub>-;
              -(CH_2)_v-C(O)-NR^{202}-(CH_2)_z-;
              -(CH_2)_v - NR^{202} - C(O) - (CH_2)_z - ;
              -(CH_2)_{v}-NR^{202}-C(O)-NR^{203}-(CH_2)_{z}-;
              -S(O)x-(CR202R203),-;
155
              - (CR202R203),-S(O),-;
              -S(O)x-(CR202R203)y-O-;
             -S(O),-(CR202R203),-C(O)-;
             -O- (CH<sub>2</sub>),-;
160
             - (CH<sub>2</sub>),-O-;
             -S-; and
             -0-;
             or R200 represents a bond:
             R^{201} represents one or more radicals selected from
       the group consisting of hydrido, halogen, hydroxy,
165
       carboxy, keto, alkyl, hydroxyalkyl, haloalkyl,
       cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,
       aralkyl, heterocyclylalkylene, alkylcarbonyl,
       hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl,
170
       haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene,
       alkoxycarbonyl, carboxyalkylcarbonyl,
       alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,
       alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl,
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1192

alkylamino, aralkylamino, alkylaminoalkylene,
aminocarbonyl, alkylcarbonylamino,

 ${\it alkylcarbonylaminoalkylcarbonyl,} \\ {\it alkylaminoalkylcarbonylamino,} \\$

aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino, alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene,

alkylimidocarbonyl, amidino, alkylamidino, aralkylamidino, guanidino, guanidinoalkylene, and alkylsulfonylamino; and

 R^{202} and R^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

y and z are independently 0, 1, 2, 3, 4, 5 or 6 wherein y + z is less than or equal to 6; and

x is 0, 1 or 2; or

 \mbox{R}^2 is -NHCR $^{204}\mbox{R}^{205}$ wherein \mbox{R}^{204} is alkylaminoalkylene, and \mbox{R}^{205} is aryl; or

190 R^2 is $-C(NR^{206})R^{207}$ wherein R^{206} is selected from hydrogen and hydroxy, and R^{207} is selected from alkyl, aryl and aralkyl; or

R2 has the formula:

$$- \begin{bmatrix} R^{30} \\ C - (CH_2) \end{bmatrix} - \begin{bmatrix} H \\ C \\ R^{34} \end{bmatrix}_{m} R^{32}$$
(TTT)

195 wherein:

185

200

j is an integer from 0 to 8; and m is 0 or 1; and

R³⁰ and R³¹ are independently selected from hydrogen, alkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, aminoalkyl, alkylaminoalkyl, aminocarbonylalkyl, alkoxyalkyl, and alkylcarbonyloxyalkyl; and

R³² is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl,

205 alkylcarbonylalkylene, arylcarbonylalkylene, and

1193

heterocyclylcarbonylaminoalkylene;

 R^{33} is selected from hydrogen, alkyl, -C(0) $R^{35},$ -C(0) $OR^{35},$ -SO $_2R^{36},$ -C(0) $NR^{37}R^{38},$ and -SO $_2NR^{39}R^{40},$ wherein

210 R³⁵, R³⁶, R³⁷, R³⁸, R³⁹ and R⁴⁰ are independently selected from hydrocarbon, heterosubstituted hydrocarbon and heterocyclyl; and

 \mathbb{R}^{34} is selected from hydrogen, alkyl, aminocarbonyl, alkylaminocarbonyl, and arylaminocarbonyl; or

 \mathbb{R}^2 is $-\mathbb{CR}^{41}\mathbb{R}^{42}$ wherein \mathbb{R}^{41} is aryl, and \mathbb{R}^{42} is hydroxy; and

R³ is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

220

215

wherein the \mathbb{R}^3 pyridinyl, pyrimidinyl, quinolinyl, purinyl groups are substituted with one or more radicals independently selected from keto, haloarylamino,

alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxyarylamino, alkylsulfonylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, alkylheterocyclylalkylamino,

230 heterocyclylheterocyclylalkylamino,
 alkoxycarbonylheterocyclylamino and haloalkylsulfonyl;
 and

wherein the R^3 maleimidyl, pyridonyl, thiazolyl, thiazolylamino,

groups may be optionally substituted with one or more

235

240

245

250

265

radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkenylamino, alkynylamino, cycloalkylamino, cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino,

heterocyclylheterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino, haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR⁴⁴R⁴⁵ wherein R⁴⁴ is alkylcarbonyl or amino, and R⁴⁵ is alkyl or aralkyl; and

R⁴ is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R⁴ is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl,

1195

alkylsulfinylalkylene, arylsulfinylalkylene,
alkylsulfonyl, alkylsulfonylalkylene,
arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy,
aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl,
alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano,
nitro, alkylamino, arylamino, alkylaminoalkylene,
arylaminoalkylene, aminoalkylamino, and hydroxy;

provided R^3 is not 2-pyridinyl when R^4 is a phenyl ring containing a 2-hydroxy substituent and when R^1 is hydrido; and

provided R3 is not

275

285

290

280 (IV) (V)

wherein \mathbb{R}^{43} is selected from hydrogen, alkyl, aminoalkyl, alkoxyalkyl, alkenoxyalkyl, and aryloxyalkyl; and

further provided R^2 is selected from aryl, heterocyclyl, unsubstituted cycloalkyl and cycloalkenyl when R^4 is hydrido; and

further provided that R4 is not methylsulfonylphenyl or aminosulfonylphenyl; and

further provided that R^1 is not methylsulfonylphenyl; or

a pharmaceutically-acceptable salt or tautomer thereof.

131. A compound of Formula IA

1196

wherein

R¹ is selected from hydroxy and alkoxyaryl; and
R² is selected from hydrido, halogen, mercapto,
alkyl, alkenyl, alkynyl, aryl, heterocyclyl, haloalkyl,
hydroxyalkyl, aralkyl, alkylheterocyclyl,
heterocyclylalkyl, heterocyclylheterocyclyl,

heterocyclylalkylheterocyclyl, alkylamino, alkenylamino, alkynylamino, arylamino, aryl(hydroxyalkyl)amino, heterocyclylamino, heterocyclylalkylamino, aralkylamino, N-alkyl-N-alkynyl-amino, aminoalkyl, aminoaryl, aminoalkylamino, aminocarbonylalkylene,

arylaminoalkylene, alkylaminoalkylene, arylaminoarylene, alkylaminoarylene, alkylaminoalkylamino, alkylcarbonylaminoalkylene, aminoalkylcarbonylaminoalkylene, alkylaminoalkylcarbonylamino, cycloalkyl, cycloalkenyl,

aminoalkylthio, alkylaminocarbonylalkylthio, alkoxy, alkylaminocarbonylalkylthio, alkoxy, heterocyclyloxy, alkylthio, cyanoalkylthio, alkenylthio, alkynylthio, carboxyalkylthio, arylthio, heterocyclylthio, alkoxycarbonylalkylthio, alkylsulfinyl,

alkylsulfonyl, carboxy, carboxyalkyl, alkoxyalkyl, alkoxyalkylthio, carboxycycloalkyl, carboxycycloalkenyl, carboxyalkylamino, alkoxycarbonyl, heterocyclylcarbonyl, alkoxycarbonylalkyl, alkoxycarbonylalkylamino, alkoxycarbonylheterocyclyl,

alkoxycarbonylheterocyclylcarbonyl, alkoxyalkylamino, alkoxycarbonylaminoalkylene, alkoxycarbonylaminoalkoxy, alkoxycarbonylaminoalkylamino, heterocyclylsulfonyl,

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aralkythio, heterocyclylalkylthio, aminoalkoxy,
      cyanoalkoxy, carboxyalkoxy, aryloxy, aralkoxy,
35
      alkenyloxy, alkynyloxy, and heterocyclylalkyloxy; wherein
      the aryl, heterocyclyl, heterocyclylalkyl, cycloalkyl and
      cycloalkenyl groups may be optionally substituted with
      one or more radicals independently selected from halo,
      keto, amino, alkyl, alkenyl, alkynyl, aryl, heterocyclyl,
40
      aralkyl, heterocyclylalkyl, epoxyalkyl,
      amino(hydroxyalkyl) carboxy, alkoxy, aryloxy, aralkoxy,
      haloalkyl, alkylamino, alkynylamino,
      alkylaminoalkylamino, heterocyclylalkylamino,
      alkylcarbonyl, alkoxycarbonyl, alkylsulfonyl,
45
      arylsulfonyl, and aralkylsulfonyl; or
             R^2 is R^{200}-heterocyclyl-R^{201}, R^{200}-aryl-R^{201}, or R^{200}-
      cycloalkyl-R201 wherein:
             R200 is selected from:
             - (CR202R203),-;
50
             -C(0)-;
             -C(0)-(CH2)v-;
             -C(O)-O-(CH<sub>2</sub>),-;
             - (CH<sub>2</sub>), - C(O) -;
             -O-(CH2),-C(O)-;
55
             -NR<sup>202</sup>-;
             -NR<sup>202</sup>-(CH<sub>2</sub>),-;
             - (CH<sub>2</sub>),-NR<sup>202</sup>-;
             - (CH<sub>2</sub>) - NR<sup>202</sup> - (CH<sub>2</sub>) - ;
             -(CH_2)_v-C(O)-NR^{202}-(CH_2)_v-;
             -(CH_2)_v-NR^{202}-C(O)-(CH_2)_z-;
60
             -(CH<sub>2</sub>)_v - NR<sup>202</sup> - C(O) - NR<sup>203</sup> - (CH<sub>2</sub>)_v - ;
            -S(O)x-(CR202R203)v-;
            -(CR^{202}R^{203})_{v}-S(0)_{x}-;
             -S(O),-(CR202R203),-O-;
            -S(O)_{x}-(CR^{202}R^{203})_{y}-C(O)-;
65
             -O-(CH<sub>2</sub>),-;
             - (CH<sub>2</sub>),-O-;
             -S-: and
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-0-;

70 or R²⁰⁰ represents a bond;

 R^{201} represents one or more radicals selected from the group consisting of hydrido, halogen, hydroxy, carboxy, keto, alkyl, hydroxyalkyl, haloalkyl, cycloalkyl, alkenyl, alkynyl, aryl, heterocyclyl,

75 aralkyl, heterocyclylalkylene, alkylcarbonyl, hydroxyalkylcarbonyl, cycloalkylcarbonyl, arylcarbonyl, haloarylcarbonyl, alkoxy, alkoxyalkylene, alkoxyarylene, alkoxycarbonyl, carboxyalkylcarbonyl,

alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl,

alkylsulfonyl, alkylsulfonylalkylene, amino, aminoalkyl, alkylamino, aralkylamino, alkylaminoalkylene, aminocarbonyl, alkylcarbonylamino, alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl, alkylaminoalkylcarbonylamino,

aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino, alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene, alkylimidocarbonyl, amidino, alkylamidino, aralkylamidino, guanidino, guanidinoalkylene, and alkylsulfonylamino; and

 R^{202} and R^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

y and z are independently 0, 1, 2, 3, 4, 5 or 6 wherein y + z is less than or equal to 6; and

x is 0, 1 or 2; or

 \mbox{R}^2 is $-\mbox{NHCR}^{204}\mbox{R}^{205}$ wherein \mbox{R}^{204} is alkylaminoalkylene, and \mbox{R}^{205} is aryl; or

 \mbox{R}^2 is $-\mbox{C(NR}^{206})\mbox{R}^{207}$ wherein \mbox{R}^{206} is selected from hydrogen and hydroxy, and \mbox{R}^{207} is selected from alkyl, aryl and aralkyl; or

100 R^2 has the formula:

90

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wherein:

115

j is an integer from 0 to 8; and m is 0 or 1: and

105 R³⁰ and R³¹ are independently selected from hydrogen, alkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, aminoalkyl, alkylaminoalkyl, aminocarbonylalkyl, alkoxyalkyl, and alkylcarbonyloxyalkyl; and

R³² is selected from hydrogen, alkyl, aralkyl,
110 heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene,
aminoalkyl, alkylaminoalkyl, arylaminoalkyl,
alkylcarbonylalkylene, arylcarbonylalkylene, and
heterocyclylcarbonylaminoalkylene;

 R^{33} is selected from hydrogen, alkyl, $-C(O)R^{35}$, $-C(O)OR^{35}$, $-SO_2R^{36}$, $-C(O)NR^{37}R^{38}$, and $-SO_2NR^{39}R^{40}$, wherein

R³⁵, R³⁶, R³⁷, R³⁸, R³⁹ and R⁴⁰ are independently selected from hydrocarbon, heterosubstituted hydrocarbon and heterocyclyl; and

120 R³⁴ is selected from hydrogen, alkyl, aminocarbonyl, alkylaminocarbonyl, and arylaminocarbonyl; or

 \mbox{R}^2 is $\mbox{-CR}^{41}\mbox{R}^{42}$ wherein \mbox{R}^{41} is aryl, and \mbox{R}^{42} is hydroxy; and

R³ is selected from pyridinyl, pyrimidinyl,
125 quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl,
thiazolylalkyl, thiazolylamino.

wherein the R³ pyridinyl, pyrimidinyl, quinolinyl, 130 purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylalkyl, thiazolylamino,

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groups may be optionally substituted with one or more
radicals independently selected from halo, keto, alkyl,
aralkyl, aralkenyl, arylheterocyclyl, carboxy,
carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio,
alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl,
aralkoxy, heterocyclylalkoxy, amino, alkylamino,
alkenylamino, alkynylamino, cycloalkylamino,
cycloalkenylamino, arylamino, haloarylamino,

cycloalkenylamino, arylamino, haloarylamino, heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl,

alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino, alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl(hydroxyalkyl)amino, alkylaminoalkylamino, alkylaminoalkylamino, alkylamino

alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino, heterocyclylheterocyclylalkylamino, alkoxycarbonylheterocyclylamino, nitro, alkylaminocarbonyl, alkylcarbonylamino,

haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl, hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR⁴⁴R⁴⁵ wherein R⁴⁴ is alkylcarbonyl or amino, and R⁴⁵ is alkyl or aralkyl; and

R⁴ is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R⁴ is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio, alkylthioalkylene, arylthioalkylene, alkylsulfinyl,

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alkylsulfinylalkylene, arylsulfinylalkylene,
alkylsulfonyl, alkylsulfonylalkylene,
arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy,
aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl,
alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano,
nitro, alkylamino, arylamino, alkylaminoalkylene,
arylaminoalkylene, aminoalkylamino, and hydroxy;
provided R³ is not 2-pyridinyl when R⁴ is a phenyl
ring containing a 2-hydroxy substituent and when R¹ is
hydrido; and

further provided R^2 is selected from aryl, heterocyclyl, unsubstituted cycloalkyl and cycloalkenyl when R^4 is hydrido; and

further provided that R^4 is not methylsulfonylphenyl or aminosulfonylphenyl; or

a pharmaceutically-acceptable salt or tautomer thereof.

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- 132. A pharmaceutical composition comprising a therapeutically-effective amount of a compound, said compound selected from the compounds of any one of Claims 1, 39, 71, 82 and 94, or a pharmaceutically acceptable salt thereof.
- 133. A method of treating a TNF mediated disorder, said method comprising treating the subject having or susceptible to such disorder with a therapeutically-effective amount of a compound, said compound selected from the compounds of any one of Claims 1, 39, 71, 82 and 94, or a pharmaceutically acceptable salt thereof.
- 134. A method of treating a p38 kinase mediated disorder, said method comprising treating the subject having or susceptible to such disorder with a therapeutically-effective amount of a compound, said compound selected from the compounds of any one of Claims

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- 5 1, 39, 71, 82 and 94, or a pharmaceutically acceptable salt thereof.
 - 135. The method of Claim 134 wherein the p38 kinase mediated disorder is selected from the group of disorders consisting of bone resorption, graft vs. host reaction, atherosclerosis, arthritis, osteoarthritis, rheumatoid arthritis, gout, psoriasis, topical inflammatory disease state, adult respiratory distress syndrome, asthma, chronic pulmonary inflammatory disease, cardiac reperfusion injury, renal reperfusion injury, thrombus, glomerulonephritis, Crohn's disease, ulcerative colitis, inflammatory bowel disease and cachexia.

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- 136. The method of Claim 134 wherein the p38 kinase mediated disorder is inflammation.
- 137. The method of Claim 134 wherein the p38 kinase mediated disorder is arthritis.
- 138. The method of Claim 134 wherein the p38 kinase mediated disorder is asthma.
- 139. A method of treating inflammation, said method comprising treating the subject having or susceptible to inflammation with a therapeutically-effective amount of a compound, said compound selected from the compounds of any one of Claims 1, 39, 71, 82 and 94, or a pharmaceutically acceptable salt thereof.
- 140. A method of treating arthritis, said method comprising treating the subject having or susceptible to arthritis with a therapeutically-effective amount of a compound, said compound selected from the compounds of any one of Claims 1, 39, 71, 82 and 94, or a pharmaceutically acceptable salt thereof.

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141. A method of preparing pyrazoles of Formula IA

wherein

arylcarbonyloxyarylene, and

heterocyclylcarbonyloxyarylene; or

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R1 is selected from hydrido, hydroxy, alkyl, 5 cycloalkyl, alkenyl, cycloalkenyl, alkynyl, aryl, heterocyclyl, cycloalkylalkylene, cycloalkenylalkylene, heterocyclylalkylene, haloalkyl, haloalkenyl, haloalkynyl, hydroxyalkyl, hydroxyalkenyl, hydroxyalkynyl, aralkyl, aralkenyl, aralkynyl, arylheterocyclyl, carboxy, carboxyalkyl, alkoxyalkyl, 10 alkenoxyalkyl, alkynoxyalkyl, aryloxyalkyl, alkoxyaryl, heterocyclyloxyalkyl, alkoxyalkoxy, mercaptoalkyl, alkylthioalkylene, alkenylthioalkylene, alkylthioalkenylene, amino, aminoalkyl, alkylamino, alkenylamino, alkynylamino, arylamino, heterocyclylamino, 15 alkylsulfinyl, alkenylsulfinyl, alkynylsulfinyl, arylsulfinyl, heterocyclylsulfinyl, alkylsulfonyl, alkenylsulfonyl, alkynylsulfonyl, arylsulfonyl, heterocyclylsulfonyl, alkylaminoalkylene, alkylsulfonylalkylene, acyl, acyloxycarbonyl, 20 alkoxycarbonylalkylene, aryloxycarbonylalkylene, heterocyclyloxycarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, heterocyclyloxycarbonylarylene, alkylcarbonylalkylene, arylcarbonylalkylene. 25 heterocyclylcarbonylalkylene, alkylcarbonylarylene, arylcarbonylarylene, heterocyclylcarbonylarylene, alkylcarbonyloxyalkylene, arylcarbonyloxyalkylene, heterocyclylcarbonyloxyalkylene, alkylcarbonyloxyarylene,

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R1 has the formula

$$- \int_{H}^{R^{25}} (CH_2) = C - N_{R^{27}}$$
(II)

wherein:

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i is an integer from 0 to 9;

35 R²⁵ is selected from hydrogen, alkyl, aralkyl, heterocyclylalkyl, alkoxyalkylene, aryloxyalkylene, aminoalkyl, alkylaminoalkyl, arylaminoalkyl, alkylcarbonylalkylene, arylcarbonylalkylene, and heterocyclylcarbonylaminoalkylene; and

40 R²⁶ is selected from hydrogen, alkyl, alkenyl, alkynyl, cycloalkylalkylene, aralkyl, alkoxycarbonylalkylene, and alkylaminoalkyl; and

R²⁷ is selected from alkyl, cycloalkyl, alkynyl, aryl, heterocyclyl, aralkyl, cycloalkylalkylene, cycloalkenylalkylene, cycloalkylarylene, cycloalkylcycloalkyl, heterocyclylalkylene, alkylarylene, alkylaralkyl, aralkylarylene, alkylheterocyclyl, alkylheterocyclylalkylene, alkylheterocyclylarylene,

aralkylheterocyclyl, alkoxyalkylene, alkoxyarylene, alkoxyaralkyl, alkoxyheterocyclyl, alkoxyalkoxyarylene,

aryloxyarylene, aralkoxyarylene,
alkoxyheterocyclylalkylene, aryloxyalkoxyarylene,
alkoxycarbonylalkylene, alkoxycarbonylheterocyclyl,
alkoxycarbonylheterocyclylcarbonylalkylene, aminoalkyl,

alkylaminoalkylene, arylaminocarbonylalkylene, alkoxyarylaminocarbonylalkylene, aminocarbonylalkylene, arylaminocarbonylalkylene, alkylaminocarbonylalkylene, arylcarbonylalkylene, alkoxycarbonylarylene, aryloxycarbonylarylene, alkylaryloxycarbonylarylene,

arylcarbonylarylene, alkylarylcarbonylarylene,
alkoxycarbonylalkoxylarylene,
alkoxycarbonylalkoxylarylene,
heterocyclylcarbonylalkylarylene, alkylthioalkylene,

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cycloalkylthioalkylene, alkylthioarylene, 65 aralkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, arylsulfonylaminoalkylene, alkylsulfonylarylene, and alkylaminosulfonylarylene; wherein said alkyl, cycloalkyl, aryl, heterocyclyl, aralkyl, heterocyclylalkylene, alkylheterocyclylarylene, alkoxyarylene, aryloxyarylene, arylaminocarbonylalkylene, 70 aryloxycarbonylarylene, arylcarbonylarylene, alkylthioarylene, heterocyclylthioarylene, arylthioalklylarylene, and alkylsulfonylarylene groups may be optionally substituted with one or more radicals independently selected from alkyl, halo, haloalkyl, 75 alkoxy, keto, amino, nitro, and cyano; or R^{27} is $-CHR^{28}R^{29}$ wherein R^{28} is alkoxycarbonyl, and R^{29}

is selected from aralkyl, aralkoxyalkylene, heterocyclylalkylene, alkylheterocyclylalkylene, alkylthioalkylene, and aralkylthioalkylene; wherein said aralkyl and heterocylcyl groups may be optionally substituted with one or more radicals independently selected from alkyl and nitro; or

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and

R²⁶ and R²⁷ together with the nitrogen atom to which they are attached form a heterocycle, wherein said heterocycle is optionally substituted with one or more radicals independently selected from alkyl, aryl, heterocyclyl, heterocyclylalkylene,

alkylheterocyclylalkylene, aryloxyalkylene, alkoxyarylene, alkylaryloxyalkylene, alkylcarbonyl, alkoxycarbonyl, aralkoxycarbonyl, alkylamino and alkoxycarbonylamino; wherein said aryl, heterocyclylalkylene and aryloxyalkylene radicals may be optionally substituted with one or more radicals independently selected from halogen, alkyl and alkoxy;

R² is selected from mercapto,
aryl(hydroxyalkyl)amino, N-alkyl-N-alkynyl-amino,

```
100
        aminocarbonylalkylene, alkylcarbonylaminoalkylene,
        aminoalkylcarbonylaminoalkylene.
        alkylaminoalkylcarbonylamino, aminoalkylthio,
        alkylaminocarbonylalkylthio,
        alkylaminoalkylaminocarbonylalkylthio, cyanoalkylthio,
        alkenylthio, alkynylthio, carboxyalkylthio,
105
        alkoxycarbonylalkylthio, alkylsulfinyl, alkylsulfonyl,
        alkoxyalkyl, alkoxyalkylthio, alkoxycarbonylalkylamino,
        alkoxycarbonylaminoalkylene, alkoxycarbonylaminoalkoxy,
        aralkythio, heterocyclylalkylthio, aminoalkoxy,
110
        cyanoalkoxy, carboxyalkoxy, aryloxy, aralkoxy,
        alkenyloxy, alkynyloxy, and heterocyclylalkyloxy; or
               R^2 is R^{200}-heterocyclyl-R^{201}, R^{200}-aryl-R^{201}, or R^{200}-
        cycloalkyl-R201 wherein:
               R200 is selected from:
115
               - (CR202R203),-;
               -C(0)-;
               -C(O)-(CH<sub>2</sub>),-;
               -C(O)-O-(CH<sub>2</sub>),-;
               - (CH<sub>2</sub>),-C(O)-;
               -O- (CH<sub>2</sub>) ,-C (O) -;
120
               -NR202-;
               -NR<sup>202</sup>- (CH<sub>2</sub>),-;
               - (CH<sub>2</sub>) v-NR<sup>202</sup>-;
               -(CH_2)_v - NR^{202} - (CH_2)_v - i
125
               - (CH<sub>2</sub>)<sub>v</sub>-C(O)-NR<sup>202</sup>-(CH<sub>2</sub>)<sub>v</sub>-;
               -(CH_2)_v - NR^{202} - C(O) - (CH_2)_z - ;
               -(CH_2)_v-NR^{202}-C(O)-NR^{203}-(CH_2)_v-i
               -S(O),-(CR202R203),-;
              -(CR^{202}R^{203})_{v}-S(O)_{x}-;
130
              -S(0) - (CR^{202}R^{203}) - O - ;
               -S(0) - (CR^{202}R^{203}) - C(0) - ;
              -O-(CH<sub>2</sub>),-;
              - (CH<sub>2</sub>)<sub>v</sub>-O-;
              -S-: and
135
              -0-;
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or R²⁰⁰ represents a bond:

 R^{201} represents one or more radicals selected from the group consisting of hydroxy, hydroxyalkyl, cycloalkyl, hydroxyalkylcarbonyl, cycloalkylcarbonyl,

- arylcarbonyl, haloarylcarbonyl, alkoxyalkylene, alkoxyarylene, carboxyalkylcarbonyl, alkoxyalkylcarbonyl, heterocyclylalkylcarbonyl, alkylsulfonylalkylene, aminoalkyl, aralkylamino, alkylaminoalkylene, aminocarbonyl, alkylcarbonylamino,
- alkylcarbonylaminoalkylene, alkylaminoalkylcarbonyl, alkylaminoalkylcarbonylamino, aminoalkylcarbonylaminoalkyl, alkoxycarbonylamino, alkoxyalkylcarbonylamino, alkoxycarbonylaminoalkylene, alkylimidocarbonyl, amidino, alkylamidino, aralkylamidino, guanidino, guanidinoalkylene, and
- alkylsulfonylamino; and

 \mbox{R}^{202} and \mbox{R}^{203} are independently selected from hydrido, alkyl, aryl and aralkyl; and

y and z are independently 0, 1, 2, 3, 4, 5 or 6 wherein y + z is less than or equal to 6; and

x is 0, 1 or 2; or

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 \mbox{R}^2 is $\mbox{-NHCR}^{204}\mbox{R}^{205}$ wherein \mbox{R}^{204} is alkylaminoalkylene, and \mbox{R}^{205} is aryl; or

 R^2 is $-C(NR^{206})R^{207}$ wherein R^{206} is selected from hydrogen and hydroxy, and R^{207} is selected from alkyl, aryl and aralkyl; and

 R^3 is selected from pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl, thiazolylamino,

wherein the R^3 pyridinyl, pyrimidinyl, quinolinyl, purinyl, maleimidyl, pyridonyl, thiazolyl,

1208

thiazolylalkyl, thiazolylamino,

$$\begin{pmatrix} 1 & 1 & 1 \\ 1 & 1 & 1 \\ 1 & 1 & 1 \end{pmatrix}$$
; and $\begin{pmatrix} 1 & 1 & 1 \\ 1 & 1 & 1 \\ 1 & 1 & 1 \end{pmatrix}$

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groups may be optionally substituted with one or more radicals independently selected from halo, keto, alkyl, aralkyl, aralkenyl, arylheterocyclyl, carboxy,

carboxyalkyl, alkoxy, aryloxy, alkylthio, arylthio, alkylsulfinyl, arylsulfinyl, alkylsulfonyl, arylsulfonyl, aralkoxy, heterocyclylalkoxy, amino, alkylamino, alkenylamino, alkynylamino, cycloalkylamino, cycloalkenylamino, arylamino, haloarylamino,

heterocyclylamino, aminocarbonyl, cyano, hydroxy, hydroxyalkyl, alkoxyalkylene, alkenoxyalkylene, aryloxyalkyl, alkoxyalkylamino, alkylaminoalkoxy, alkoxycarbonyl, aryloxycarbonyl, heterocyclyloxycarbonyl, alkoxycarbonylamino, alkoxyarylamino, alkoxyaralkylamino, aminosulfinyl, aminosulfonyl, alkylsulfonylamino,

alkylaminoalkylamino, hydroxyalkylamino, aralkylamino, aryl (hydroxyalkyl) amino, alkylaminoalkylaminoalkylamino, alkylheterocyclylamino, heterocyclylalkylamino, alkylheterocyclylalkylamino, aralkylheterocyclylamino, heterocyclylalkylamino, heterocyclylalkylamino,

heterocyclylheterocyclylalkylamino,
 alkoxycarbonylheterocyclylamino, nitro,
 alkylaminocarbonyl, alkylcarbonylamino,
 haloalkylsulfonyl, aminoalkyl, haloalkyl, alkylcarbonyl,
 hydrazinyl, alkylhydrazinyl, arylhydrazinyl, and -NR⁴⁴R⁴⁵
 wherein R⁴⁴ is alkylcarbonyl or amino, and R⁴⁵ is alkyl or
 aralkyl; and

 R^4 is selected from hydrido, alkyl, alkenyl, alkynyl, cycloalkyl, cycloalkenyl, aryl, and heterocyclyl, wherein R^4 is optionally substituted with one or more radicals independently selected from halo, alkyl, alkenyl, alkynyl, aryl, heterocyclyl, alkylthio, arylthio,

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alkylthioalkylene, arylthioalkylene, alkylsulfinyl, alkylsulfinylalkylene, arylsulfinylalkylene, alkylsulfonyl, alkylsulfonylalkylene, arylsulfonylalkylene, alkoxy, aryloxy, aralkoxy, aminocarbonyl, alkylaminocarbonyl, arylaminocarbonyl, alkoxycarbonyl, aryloxycarbonyl, haloalkyl, amino, cyano, nitro, alkylamino, arylamino, alkylaminoalkylene,

said method comprising the steps of treating a substituted ketone with an acyl hydrazide to give the pyrazole.

- 142. The process of Claim 141 wherein the process is carried out in an acidic solvent.
- 143. The process of Claim 141 wherein the acidic solvent is acetic acid.
- 144. The process of Claim 141 wherein the acidic solvent is an organic solvent containing an acid.

145. The compound:

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or a tautomer or pharmaceutically acceptable salt thereof,

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146. A compound of Claim 71 that is:

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or a tautomer or pharmaceutically acceptable salt thereof.

147. A compound of Claim 39 that is:

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or a tautomer or pharmaceutically acceptable salt thereof.

148. The compound:

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or a tautomer or pharmaceutically acceptable salt thereof.

149. A compound of Claim 1 that is:

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or a tautomer or pharmaceutically acceptable salt thereof.

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150. The compound:

or a tautomer or pharmaceutically acceptable salt thereof.

30 151. A compound of Claim 1 that is:

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or a tautomer or pharmaceutically acceptable salt thereof.

152. A compound of Claim 1 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

153. A compound of Claim 1 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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154. A compound of Claim 39 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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155. A compound of Claim 1 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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156. A compound of Claim 82 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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157. A compound of Claim 42 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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158. A compound of Claim 71 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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159. A compound of Claim 71 that is:

or a tautomer or pharmaceutically acceptable salt thereof.

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160. A compound of Claim 70 wherein R^{404a} is metachloro or para-chloro.

Internation pplication No PCT/US 99/26007

A. CLASSIFICATION OF SUBJECT MATTER
IPC 7 C070401/04 A61k A61K31/415 A61K31/47 A61K31/445 A61K31/44 A61K31/50 A61K31/505 A61K31/52 CO7D405/14 C070401/14 C070409/14 C07D403/04 C07D487/04 C07D473/00 C070413/14

According to International Patent Classification (IPC) or to both national classification and IPC

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

IPC 7 CO7D A61K A61P

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Escronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category *	Citation of document, with Indication, where appropriate, of the relevant passages	Relevant to claim No.
X	WO 96 03385 A (SEARLE & CO ; LEE LEN F (US); PENNING THOMAS D (US); KRAMER STEVEN) 8 February 1996 (1996-02-08) cited in the application abstract; claims 1,8-10; examples 1-15 page 9 -page 73	1,39,71, 82,93, 94,101, 126-140
A	EP 0 846 687 A (LILLY CO ELI) 10 June 1998 (1998-06-10) abstract; examples page 21; table 1A page 23 -page 25; table 2A -/	1,39,71, 82,93, 94,10i

X	Further documents are listed in the	continuation of box C.
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Patent family members are listed in annex.

- Special categories of cited documents:
- "A" document defining the general state of the art which is not considered to be of particular relevence
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- "&" document member of the same patent family

Paisdor, B

Date of the actual completion of the international search Date of mailing of the international search report 6 April 2000 18/04/2000 Name and mailing address of the ISA Authorized officer European Patent Office, P.B. 5816 Patentiaan 2 NL – 2280 HV Rijewijk Tal. (+31-70) 340-2040, Tx. 31 651 epo ni, Fax: (+31-70) 340-3016

Internation. .pplication No PCT/US 99/26007 A. CLASSIFICATION OF SUBJECT MATTER IPC 7 C07D417/14 C07D C07D471/04 //(C07D487/04,293:00, A61P29/00 231:00),(C07D471/04,221:00,209:00) According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where prectical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriats, of the relevant passages Relevant to claim No. A EP 0 846 686 A (PFIZER LTD ; PFIZER (US)) 1,39,71, 10 June 1998 (1998-06-10) 82,93, 94,101 abstract; claims 1,15 page 19; example A24 WO 94 19350 A (OKU TERUO ; KAWAI YOSHIO A 1,39,71. (JP); TANAKA HIROKAZU (JP); FUJISAWA 82.93. PHARM) 1 September 1994 (1994-09-01) 94,101 page 53; example 8 A EP 0 531 901 A (FUJISAWA PHARMACEUTICAL 1,39,71. CO) 17 March 1993 (1993-03-17) 82,93, 94,101 abstract pages 49 - 51, preparations page 52; example 1 -/--Further documents are listed in the continuation of box C. X Patent family members are listed in annex. Special categories of cited documente : T* later document published effer the international filing date or priority date and not in conflict with the application but clied to understand the principle or theory underlying the "A" document defining the general state of the art which is not considered to be of particular relevance Invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date ." document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such document. "O" document referring to an oral disclosure, use, exhibition or other means ments, such combination being obvious to a person skilled in the art. *P" document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the International search report 6 April 2000 Name and mailing address of the ISA

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Internation. aplication No PCT/IIS 99/26007

0./04=41:-		PCT/US 99/26007
C.(Continue	MION) DOCUMENTS CONSIDERED TO BE RELEVANT Citation of document, with indication, where appropriats, of the relevant passages	
Category	Chamon of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	DD 295 374 A (STERLING DRUG 1NC) 31 October 1991 (1991-10-31)	1,39,71, 82,93,
	page 7 -page 8; claims; examples	94,101
A	WO 95 31451 A (SMITHKLINE BEECHAM CORP; ADAMS JERRY LEROY (US); GALLAGHER TIMOTHY) 23 November 1995 (1995-11-23)	1,39,71, 82,93, 94,101,
	page 1 -page 3; claim 1 page 16 -page 19; examples	126-140
Ρ,Χ	WO 98 52937 A (ANANTANARAYAN ASHOK ;STEALEY M1CHAEL A (US); CLARE M1CHAEL (US); G) 26 November 1998 (1998-11-26) abstract; claims page 35 -page 49; examples	1-160
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Interna vonal application No.

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Box I	Observations where certain claims were found unsearchable (Continuation of Item 1 of first sheet)
This inte	emational Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
1. X	Claims Nos.: 133-140 because they relate to subject matter not required to be searched by this Authority, namely: Remark: Although claims 133-140 are directed to a method of treatment of the human/animal body, the search has been carried out and based on the alleged
2.	effects of the compound/composition. Claims Nos.: because they relate to parts of the International Application that do not comply with the prescribed requirements to such an extent that no meaningful International Search can be carried out, specifically:
3.	Claims Noa.: because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Box II	Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
PHO man	emational Searching Authority tound multiple inventions in this international application, as follows:
1.	As all required additional search fees were timely paid by the applicant, this International Search Report covers all searchable claims.
2.	As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite payment of any additional fee.
з	As only some of the required additional search fees were timely paid by the applicant, this international Search Report covers only those claims for which fees were paid, specifically claims Nos.:
4.	No required additional search fees were timely paid by the applicant. Consequently, this International Search Report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
Remark c	The additional search fees were accompanied by the applicant's protest. No protest accompanied the payment of additional search fees.

WILLIAM DEALCH REPORT

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